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(54) Title: COMPOSITIONS THAT INHIBIT PROLIFERATION OF CANCER CELLS

(57) Abstract: Disclosed are compositions and methods for reducing the proliferation of cancer cells through targeted interactions with integrins.

COMPOSITIONS THAT INHIBIT PROLIFERATION OF CANCER CELLS

This application claims priority to United States Provisional Application No. 60/239,705 filed on October 12, 2000, entitled "Agents promoting apoptosis in cancer cells via interruption of oncogene-induced integrin signaling," which application is herein incorporated by reference in its entirety and to United States Provisional Application No. 60/242,812 filed on October 24, 2000 entitled, "Agents promoting apoptosis in cancer cells via interruption of oncogene-induced integrin signaling", and which application is herein incorporated by reference in its entirety.

I. BACKGROUND OF THE INVENTION

In cancer cells multiple oncogenic lesions cooperate in malignant transformation. Such cooperation permits survival and proliferation of tumor cells in absence of contact with extracellular matrix (ECM), suggesting that tumor cell survival and proliferation have become independent of the engagement of integrin signaling by ECM.

Carcinogenesis is caused by multiple cooperating genetic lesions leading to a progressive deregulation of cellular signaling and cell cycle restriction point control. The mutations involved result in oncogene activation or loss of tumor-suppressor gene function. Typically, single oncogenes are insufficient to cause malignant transformation because they simultaneously induce signals stimulating and inhibiting cell growth. As a result cell proliferation remains restricted. In contrast, cooperating oncogenic lesions act in concert to disable such inhibitory signals while reinforcing the growth-promoting stimuli. The co-operation of oncogenic lesions involves integration of multiple signals converging on the regulation of cell cycle-dependent kinase complexes (Lloyd et al., 1997; Perez-Roger et al., 1999; Roper et al., 2001; Sewing et al., 1997).

Disclosed herein are compositions and methods that show survival of various transformed cell types requires cell-autonomous (autocrine) integrin signaling activity. This activity is induced by cooperating oncogenic lesions and involves induction of integrin receptor and ligand components such as integrin alpha6 and integrin beta4, and laminin5-gamma2 chains. Blocking of integrin or the laminin ligand function induces rapid apoptosis of the transformed cells, even when growing in presence of ECM. In contrast, normal cells remain viable when exposed to the same treatment.

The disclosed compositions and methods are related to the cooperation of oncogenic lesions controlling the ability of transformed cells to proliferate in the absence of contact with the extra-cellular matrix (ECM). As taught herein, oncogenes cooperate to promote a cell-autonomous (autocrine) integrin signaling loop that proves essential for the survival of various transformed cell

types. As this signaling loop is not established in corresponding normal cells, the signaling components of this loop constitute attractive targets for cancer therapy.

II. SUMMARY OF THE INVENTION

In accordance with the purposes of this invention, as embodied and broadly described berein, this invention, in one aspect, relates to compositions and methods related to integrin mediated cancer cell growth.

Additional advantages of the invention will be set forth in part in the description which follows, and in part will be obvious from the description, or may be learned by practice of the invention. The advantages of the invention will be realized and attained by means of the elements and combinations particularly pointed out in the appended claims. It is to be understood that both the foregoing general description and the following detailed description are exemplary and explanatory only and are not restrictive of the invention, as claimed.

III. BRIEF DESCRIPTION OF THE DRAWINGS

The accompanying drawings, which are incorporated in and constitute a part of this

15 specification, illustrate several embodiments of the invention and together with the description, serve to explain the principles of the invention.

Figure 1 shows a series of schematics representative of the disclosed relationships and compositions. In normal cells integrin receptors signal to suppress programmed cell death (apoptosis) when engaged by appropriate extra-cellular matrix (ECM) ligands. When receptor-20 ligand interaction is lost, the cells undergo apoptosis due to the lack of survival signals (Figure 1-. Panel 1). In cancer cells multiple oncogenic lesions cooperate to cause malignant transformation. Such cooperation permits survival and proliferation of tumor cells independent of integrin receptor-ECM interactions. This property has been termed anchorage-independence (Panel 2). We have discovered the mechanistic basis of anchorage-independence. The transformed cells 25 replace the requirement for ECM-dependent signaling with a surrogate integrin signaling loop on which they rely for survival. In colonic epithelial cells, activation of Ras in combination with APC (adenomatous polyposis coli) or p53 mutations leads to induction of integrin receptor and ligand components such as integrin alpha6 and laminin gamma2 chains. As a consequence, laminindependent activation of alpha6/beta4 integrin receptors signals to inhibit caspase activity and thus 30 to suppress apoptosis (Panel 3). Ablation of integrin alpha6, laminin ligand function or alpha6/beta4 integrin receptor function induces apoptosis of the transformed cells, even when growing in the presence of ECM. In contrast, non-transformed control cells remain viable when exposed to the same treatment, indicating that the dependence of the transformed cells on autocrine integrin signaling may be a particular feature of the cancer cell phenotype. The essential 35 role of alpha6 integrin extends to various transformed cell types including mesenchymal and

SW480 human colon carcinoma cells. Thus inhibition of laminin/alpha6 integrin-mediated signaling is an important method to induce cancer cell-specific death in a variety of cell types (Panel 4).

Figure 2 shows that oncogene cooperation protects cells from apoptosis. Figure 2A shows 5 control, APCm, Ras, APCm+Ras cells that were detached from collagen IV substrates with Trypsin/EDTA and kept in suspension at 2x10⁵ cells/ml in RPMI 10%FCS for 12 h at 39°C. Subsequently TUNEL analysis was performed on poly-lysine treated slides. The percentage of tunel-positive cells was determined by immunofluorescence microscopy. Figure 2B shows control, APCm, Ras, APCm Ras, SW480 cells were detached and maintained in suspension as 10 described in (A). SW480 beta4 dn/Gal4VPER cells, express a 40H-tamoxifen-inducible dominant-negative form of the beta4 integrin. Cells were pelleted and protein extracts were prepared in 300 µl of 50 mM Tris-HCl, pH 7.4; 1% NP40; 0.25% sodium deoxycholate; 150 mM NaCl; 1 mM EGTA. 200 μl of extracts were incubated for 10 min with 2 μl of Caspase 3 fluorometric Substrate (Upstate Biotechnology). Vmax of caspase activity was determined by 15 measuring the fluorescence at 460 nm after excitation at 380 nm for 1 h. Figure 2C shows laminin gammal and gamma2-specific peptides were added to APCmin+Ras cells at the concentration of 100µg/ml. Caspase activity was measured as in (B). Figure 2D shows caspase 8 and caspase 9 activities were measured as in (B) using caspase 8 and caspase 9-specific fluorimetric substrates (Upstate Biotechnologies).

Figure 3 shows alterations of integrin and laminin expression profiles in malignant cell transformation. Figure 3A shows integrin expression and figure 3B shows laminin expression. The indicated cell populations were cultured on collagen IV-coated dishes at 39°C in RPMI 10% FCS. Total RNA was extracted from 10⁶ cells for each sample and used for RT-PCR (laminins, alpha4, gamma2 and GAPDH) or RNase protection. For RT-PCR, cDNA was subjected to 28 cycles (linear range) of PCR amplification. PCR products were analyzed on a 2% agarose gel. For RNase protection, 10 μg of total RNA was used per reaction. Products were resolved on a 4.5% polyacrylamide/10M urea gel.

Figure 4 shows alpha6/beta4 integrin is engaged by the laminin gamma2 chain to activate Shc (a src homology domain containing protein). Figure 4A shows cells detached from collagen IV coated dishes with 3mM EDTA (in PBS) were incubated on ice with the indicated antibodies for 1 h. Cells were then plated on 96well dishes coated with gamma1 or gamma2-specific peptides and were permitted to attach for 30 min in RPMI medium at 39°C. After incubation, wells were washed with RPMI. The percentage of attached cells per well was measured by hexoaminidase activity after lysis of the cells in the well and incubation with the substrate p-nitrophenol-N-acetyl-35 β-D-glucosaminide for 5 h at 37°C. Figure 4B shows cells indicated were detached as in (A) and

resuspended at 10⁷ cells/ml in PBS. After 1 h of incubation on ice, cells were pelleted by centrifugation 5 min at 900 rpm and resuspended in RPMI containing the phosphatase inhibitor pervanadate at 10µM. The cells were incubated with beta1 and beta4 integrin antibodies as well as the laminin gamma2-specific peptide for 40 min. Protein extracts were prepared and subjected to an immuno-precipitation with an anti-Shc antibody. Phospho-tyrosine was detected with a phopho-tyrosine-specific antibody in IP-western-blots (upper panels). The levels of Shc protein were monitored with a Shc-specific antibody reusing the same membrane (lower panels).

Figure 5 shows integrin alpha6/beta4 and Laminin gamma2 chain expression is essential for survival of transformed cells. Figure 5A1 shows the indicated cell populations were infected at 10 an MOI of 2 with recombinant retroviruses expressing anti-sense RNA for alpha6 integrin, the gamma2 laminin chain or a beta4 integrin dominant-negative mutant together with a puromycin resistance gene. After 2 weeks of selection with puromycin, colonies were stained with Giemsa and counted. APCm+Ras alpha6 and APCm+Ras p35 cells express the alpha6 integrin chain and the anti-apoptotic baculovirus p35, respectively. APCm+Ras cells were also plated on dishes pre-15 coated with a laminin gamma2-specific peptide. Figure 5A2 shows APCm+Ras hygro, APCm+Ras Bcl2 and APCm+Ras p35 cells express the hygromycin resistance marker, and the anti-apoptotic proteins Bol2 or p35, respectively. The cells were infected at an MOI of 2 with retroviruses carrying the beta4 integrin dominant-negative mutant and a puromycin resistance gene, or the puromycin resistance gene alone. Cells (105) were maintained in soft agar at 39°C for 20 two weeks after which macroscopically visible colonies were counted. Figure 5A3 shows the indicated cell populations were infected at an MOI of 2 with recombinant retroviruses, drugselected and maintained as described in A1. Figure 5A4 shows the indicated cell populations were infected at an MOI of 2 with recombinant retroviruses expressing anti-sense RNA for alpha6 integrin, the beta4 integrin dominant-negative mutant or the puromycin resistance marker. Cells 25 (10⁵) were maintained in soft agar and evaluated as described in A2. Figure 5B shows the effect of the alpha6 anti-sense RNA expression on the cell surface expression of alpha6 integrin was monitored by FACS analysis using an anti-alpha6 rat monoclonal antibody and a FITC conjugated goat anti-rat antiserum as the secondary antibody (upper panel). Expression of the beta4 dominant-negative mutant was confirmed by RNAse protection. A probe overlapping the cterminal end of the beta4 dominant negative mRNA was used to measure beta4 integrin and beta4 dn expression in the same sample (lower panel). Figure 5C shows arf null mouse embryo fibroblasts (MEFs) that were infected with retroviruses expressing the oncogenes Myc and Ras. Cells were then additionally infected with a retroviruses expressing anti-sense RNA for alpha6 integrin or the beta4 integrin dominant-negative mutant as shown in (A). MEFS 35 Ras/Myc/Arfnull+alpha6 sense express the alpha6 integrin chain. Figure 4D shows SW480 human

colon carcinoma cells that were infected as described in (A) using VSV pseudo-typed viruses.

Infected cells were selected with puromycin in soft agar. Clones were counted 2 weeks after selection. SW480 beta4 dn/Gal4VPER cells, express a 4OH-tamoxifen-inducible dominant-negative form of the beta4 integrin. The cells (10⁵) were maintained in soft agar for 2 weeks in presence or absence of 4OH-tamoxifen.

IV. DETAILED DESCRIPTION

The present invention may be understood more readily by reference to the following detailed description of preferred embodiments of the invention and the Examples included therein and to the Figures and their previous and following description.

Before the present compounds, compositions, articles, devices, and/or methods are disclosed and described, it is to be understood that this invention is not limited to specific synthetic methods, specific recombinant biotechnology methods unless otherwise specified, or to particular reagents unless otherwise specified, as such may, of course, vary. It is also to be understood that the terminology used herein is for the purpose of describing particular embodiments only and is not intended to be limiting.

A. Definitions

As used in the specification and the appended claims, the singular forms "a," "an" and "the" include plural referents unless the context clearly dictates otherwise. Thus, for example, reference to "a pharmaceutical carrier" includes mixtures of two or more such carriers, and the like.

Ranges may be expressed herein as from "about" one particular value, and/or to "about" another particular value. When such a range is expressed, another embodiment includes from the one particular value and/or to the other particular value. Similarly, when values are expressed as approximations, by use of the antecedent "about," it will be understood that the particular value forms another embodiment. It will be further understood that the endpoints of each of the ranges are significant both in relation to the other endpoint, and independently of the other endpoint.

In this specification and in the claims which follow, reference will be made to a number of terms which shall be defined to have the following meanings:

"Optional" or "optionally" means that the subsequently described event or circumstance
may or may not occur, and that the description includes instances where said event or circumstance
occurs and instances where it does not.

"Primers" are a subset of probes which are capable of supporting some type of enzymatic manipulation and which can hybridize with a target nucleic acid such that the enzymatic manipulation can occur. A primer can be made from any combination of nucleotides or nucleotide

derivatives or analogs available in the art which do not interfere with the enzymatic manipulation.

"Probes" are molecules capable of interacting with a target nucleic acid, typically in a sequence specific manner, for example through hybridization. The hybridization of nucleic acids is well understood in the art and discussed herein. Typically a probe can be made from any combination of nucleotides or nucleotide derivatives or analogs available in the art.

B. Compositions and methods

Disclosed are compositions and methods related to integrins and integrin signaling. It is shown herein that integrin alpha6, integrin beta4, and laminin5, through at least up regulation of the beta and gamma chains of laminin5, are upregulated in cancer cells. Integrin alpha6 and 10 integrin beta4 interact to form the integrin receptor, A6B4. The integrin receptor A6B4 specifically interacts with laminin5 and laminin5 specifically interacts with A6B4. Furthermore, it is shown that interference with the production or function of alpha6, beta4, or the laminin gamma2 chain not only prevents proliferation of the cancer cells, dependent on the upregulation of alpha6, beta4, and the laminin gamma2 chain, but that this kills the cancer cells as well. Alpha6 and beta4 15 signaling occur through the integrin receptor A6B4. Thus, interference with the formation of A6B4, will interfere with the function of A6B4, for example, the signaling of A6B4. Thus, disclosed are compositions and methods that interfere with the function of alpha6, beta4, laminin5, the laminin gamma2 chain, or A6B4. Also disclosed are compositions and methods that interfere with the function of molecules involved with the signal transduction that is connected to either 20 alpha6, beta4, laminin5, the laminin gamma2 chain, or A6B4. Also disclosed are methods for reducing the proliferation of cancer cells, as well as methods of killing cancer cells that involve using the compositions disclosed herein that interfere, reduce, or eliminate the function or the alpha6, beta4, laminin5, the laminin gamma2 chain, or A6B4 function.

Disclosed herein is a relationship between two types of molecules in a cancer cell. The

first type of molecule is an integrin receptor, composed of integrins, and the second type of
molecule is a ligand that interacts with the integrin receptor, through the integrins. There are
specificities that exist between the integrin receptors and their ligands. One aspect, disclosed
herein is that when a cell goes from a non-cancerous state to a cancerous state, there is a coupregulation of both the ligand (or parts of the ligand, such as subunits) and the cognate integrin
receptor. The co-upregulation of both types of molecules creates an autocrine loop situation,
wherein the signaling pathways controlled by the integrin receptor become autonomously
activated, rather than exogenously activated, as would normally occur. The upregulation of both
types of molecules creates a more fully transformed cellular phenotype in which cancer cell
survival depends on the autocrine loop. Now therapeutic activities can target both points in the
autocrine loop. Specific examples, of this co-upregulation in cancer cells are disclosed herein. For

example, laminin 5 (both the beta and gamma2 chains are upregulated) and integrin receptor alpha6beta4; laminin 10/11 and the integrin receptors alpha6beta1 and alpha3beta1.

As the Examples herein indicate which integrins, which integrin receptor, and which integrin ligand are involved in conferring cancer cells ability to grow in the absence of extra cellular matrix (ECM), also disclosed are methods using these integrins, integrin receptors and ligands to identify molecules that interact with them and/or interfere with their function.

1. Compositions

Disclosed are the components to be used to prepare the disclosed compositions as well as the compositions themselves to be used within the methods disclosed herein. These and other 10 materials are disclosed herein, and it is understood that when combinations, subsets, interactions, groups, etc. of these materials are disclosed, while specific reference of each various individual and collective permutation of these compounds may not be explicitly disclosed, each is specifically contemplated and described herein. For example, if a particular beta4 or alpha6 is disclosed and discussed and a number of modifications that can be made to a number of molecules 15 including the modifications to beta4 or alpha6 are discussed, specifically contemplated is each and every combination and permutation of these modifications and the modifications that are possible unless specifically indicated to the contrary. Thus, if a class of molecules A, B, and C are disclosed as well as a class of molecules D, E, and F and an example of a combination molecule, A-D is disclosed, then even if each is not individually recited, each is individually and collectively 20 contemplated. Thus, combinations, A-E, A-F, B-D, B-E, B-F, C-D, C-E, and C-F are considered disclosed. Likewise, any subset or combination of these is also disclosed. Thus, for example, the sub-group of A-E, B-F, and C-E would be considered disclosed. This concept applies to all aspects of this application including, but not limited to, steps in methods of making and using the disclosed compositions. Thus, if there are a variety of additional steps that can be performed it is 25 understood that each of these additional steps can be performed with any specific embodiment or combination of embodiments of the disclosed methods.

a) Integrins and their ligands

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For proper embryonic development, tissue homeostasis, and wound healing, cell proliferation must be tightly regulated, both in space and over time. In particular, a cell must be able to sense its relationship to other cells and the extra-cellular matrix (ECM) and convert these positional cues into biochemical signals affecting the regulation of proliferation. Because of their ability to couple the recognition of positional cues to the activation of intracellular signaling pathways, adhesion receptors, such as integrins and cadherins, are likely to be necessary to achieve this goal.

The integrins mediate cell adhesion primarily by binding to distinct, although overlapping,

subsets of ECM proteins. Normal cells require contact with serum-derived ECM components for proliferation, differentiation and survival (e.g. Clark and Brugge, 1995; Lin and Bissell, 1993; Parise et al., 2000), a phenomenon called anchorage-dependence. This involves signaling through integrin receptors (Hynes, 1992). Fibronectin and laminin as well as other ECM proteins are known to act as ligands for integrin receptors (Akiyama et al., 1990). Integrins are transmembrane proteins forming alpha-beta chain heterodimers. Alpha and beta chain integrins are members of distinct gene families. The ligand binding specificity of the hetero dimers is determined by specific combinations of alpha and beta chain gene family members (Hynes, 1992). Ligand binding triggers signaling of integrin receptors through the cytoplasmic tail of the beta chain via interaction with various signaling components.

Integrins activate common as well as subgroup-specific signaling pathways (Clark and Brugge, 1995; Giancotti and Ruoslahti, 1999). In particular, while most integrins activate focal adhesion kinase (FAK), the α1β1, α5β1, ανβ3 and α6β4 integrins are coupled to the Rasextracellular signal-regulated kinase (ERK) signaling pathway by Shc (Mainiero et al., 1997; 15 Mainiero et al., 1995; Wary et al., 1996). She is an SH2-PTB domain adapter protein expressed in three forms, p46, p52 and p66, two of which (p46 and p52) link various tyrosine kinases to Ras by recruiting the Grb2/SOS complex to the plasma membrane (Pawson and Scott, 1997). Upon activation by SOS, Ras stimulates a kinase cascade culminating in the activation of the mitogenactivated protein kinase (MAPK) ERK (Marshall, 1995). ERK phosphorylates ternary complex 20 transcription factors, such as Elk-1 and Sap-1/2, and promotes transcription of the immediate-early gene Fos (Treisman, 1996). In primary endothelial cells and keratinocytes, mitogens and Shclinked integrins cooperate, in a synergic fashion, to promote transcription from the Fos promoter. Accordingly, ligation of integrins linked to Shc enables these cells to progress through G1 in response to mitogens, whereas ligation of other integrins results in growth arrest, even in the 25 presence of mitogens (Mainiero et al., 1997; Wary et al., 1996). She is like a binary switch controlling cell cycle progression in response to the ECM. Moreover, integrin receptors have been shown to induce intracellular signaling leading to AKT activation supporting cell survival (Lee and Juliano, 2000).

In contrast with normal cells, cancer cells generally are able to survive and proliferate in the absence of anchorage to ECM (Giancotti and Mainiero, 1994), suggesting that tumor cell survival and proliferation have become independent of the engagement of integrin signaling through ECM.

Proliferation in the absence of anchorage to ECM of secondary rat embryo fibroblasts requires the cooperation of Ras and Myc or Ras and adenovirus E1a oncogenes (Land et al., 1983; Ruley, 1983). Similarly, murine colonic epithelial cells require both activated Ras and mutation of

the adenomous polyposis coli gene (APCmin) (D'Abaco et al., 1996) in order to proliferate in suspension.

Integrins are a large family of cell surface receptor molecules that function to mediate interactions between cells and between cells and the extracellular matrix. Integrin receptors are heterodimers composed of two subunits, an alpha integrin and a beta integrin. The heterodimer forms, is expressed on the cell surface, and acts to transmit signals obtained from interactions with the extracellular matrix or other cells, through the cellular membrane and into the cytosol of the cell. The signal transduction that takes place occurs because of ligand interactions with the receptor. Integrin receptors can have a number of ligands, including collagens, fibronectins, and laminins.

There are currently at least 18 different alpha integrins and at least 8 different beta integrins that have been shown to form at least 24 different alphabeta heterodimers. Certain integrins, such as beta1, interact in a number of different heterodimers, but many subunits only form a single heterodimer, either because of structural constraints on their interactions, or cellular expression patterns that provide only a limited number of potential dimer partners. Disclosed herein are specific relationships that occur between a subset of integrins, integrin receptors, and their ligands. The disclosed relationships, revolve around the alpha6beta4 receptor, formed by the alpha6 and beta4 integrins. Of particular interest is the relationship between the ligand for the alpha6beta4 receptor, laminin5. The laminins are made up of 3 chains, an alpha chain, a beta chain, and a gamma chain. The specificity of the interaction between laminin5 and alpha6beta4 receptor is controlled by the gamma chain. Laminin5 contains a gamma2 chain which only interacts with the alpha6beta4 integrin receptor.

Integrin alpha6 has seven amino-terminal repeating segments that may fold into a seven unit beta-propeller, five n-terminal FG-GAP domains and three divalent cation sites. The transmembrane domain is followed by a short cytoplasmic tail, that is alternatively spliced in A and B forms. The alpha6 integrin chain also shows alternative splicing between repeat units III and IV, resulting in the presence or absence of Exon X2. Integrin alpha6 is processed into a heavy and a light chain that are disulphide linked. A representative allele of the human alpha6 cDNA is set forth in SEQ ID NO:1. It is understood that the disclosed functional domains as well as the others contained within alpha6 are considered separately disclosed as discreet fragments of the alpha6 protein as well as the nucleic acid that encodes them.

Integrin beta4 contains a MIDAS-like motif and four cysteine-rich repeats, three EGF-like domains in the N-terminal extracellular domain, a trans-membrane region and a long cytoplasmic tail containing two pairs of fibronectin Type III repeats. The latter are connected by a variable segment that may undergo alternative splicing. Integrin beta4 also undergoes proteolytic

processing in its cytoplasmic tail, causing the 200kD mature form to be converted to 165 and 130kD fragments. A representative allele of the human beta4 cDNA is set forth in SEQ ID NO:5. It is understood that the disclosed functional domains as well as the others contained within beta4 are considered separately disclosed as discreet fragments of the beta4 protein as well as the nucleic 5 acid that encodes them.

Laminin5 is composed of the laminin chains alpha3, beta3 and gamma2. Laminin5 can may contain either the shorter laminin alpha3A chain or the longer alpha3B chain. Laminin5 can also be trimmed by proteolytic processing of the N-terminal portion of its alpha3A chain and the N-terminal portion of the gamma2 chain. The Laminin gamma2 chain contains at least six laminin EGF-like domains (Domains III and V) with an embedded laminin B domain (Domain IV) within the N-terminal half. The c-terminal tail contains a coiled-coil domain. The N-terminal processed portion of the gamma2 chain is sufficient to bind to and activate the integrin alpha6/beta4 receptor. A representative allele of the human laminin5-gamma2 cDNA is set forth in SEQ ID NO:13. It is understood that the disclosed functional domains as well as the others contained within the laminin5 protein and laminin5-gamma2 protein are considered separately disclosed as discreet fragments of the laminin5 protein and laminin5-gamma2 protein as well as and the nucleic acid that encodes them.

Disclosed are compositions and methods for inhibiting integrin signaling, for example, integrin signaling dependent on alpha6 and beta4 integrins. For example, compositions and methods that inhibit integrin receptor signaling from, for example, the alpha6beta4 integrin receptor are disclosed. It is also understood that the integrin receptor signaling can be affected by, for example, interfering with a molecule, such as a ligand for the integrin receptor, or a downstream signaling molecule of the integrin receptor in a way that prevents the integrin receptor signal from being fully propagated. It is understood that the compositions and methods for inhibition of integrin signaling and function can be any composition or method that ultimately inhibits the cell proliferation in which the integrin is expressed, by for example killing the cell. It is understood that the compositions and methods typically can fall into three basic non-limiting classes of function regulators, which are discussed herein.

(1) Classes

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(a) Production regulators

Production regulators is a broad class of integrin function regulators that are directed at the production of the target integrin, by for example, preventing mRNA synthesis or expression of the target integrin, or by causing mRNA degradation of the target integrin which inhibits the translation of the target integrin. While production regulators, can be any type of molecule targeting any point in the integrin production pathway, typically these types of compositions will

target either the mRNA expression or the protein expression of the integrin. For example, if beta4 integrin, alpha6 integrin, or the gamma2 subunit of laminin5, which has been shown herein to be upregulated in cancer cells and which causes the cancer cell to be able to live in the absence of the ECM, was the target integrin, a typical production regulator of beta4 integrin, alpha6 integrin, or the gamma2 subunit of laminin5 would be for example, an antisense molecule that targeted the mRNA of beta4 integrin, alpha6 integrin, or the gamma2 subunit of laminin5. It is also understood that a production regulator could also target any molecule that is disclosed herein, or is within the signaling chain associated with a target integrin or integrin receptor. For example, the inhibition of the production of the ligand for a target integrin receptor is one way of inhibiting integrin function of that receptor. Thus, production regulators can either inhibit or enhance integrin production.

(b) Integrin to integrin regulators

Another type of integrin function regulator is an integrin-integrin regulator. This type of function regulator, typically prevents integrins from interacting to form a functional integrin receptor. For example, an integrin to integrin regulator could be a composition that would interact with beta4 in a way that would prevent beta4 from interacting with alpha6 to form the alpha6beta4 integrin receptor or it could be a composition that would interact with alpha6 in a way that would prevent alpha6 from interacting with beta4 to form the alpha6beta4 integrin receptor. It is also contemplated that the function regulators of the integrin to integrin interaction can affect the signaling pathways dependent on integrin receptors containing alph6 or beta4 integrins. It is not required that an integrin to integrin regulator prevent an integrin from interacting with all of the possible integrin partners it could interact with, just that it prevent the interaction of the target integrin with another specific integrin. For example, the integrin alpha6 interacts with beta1 integrin and beta4 integrin. In certain embodiments, the compositions interfere with alpha6-beta4 interactions but do not interfere with alpha6-beta1 interactions.

(c) Integrin to other molecule regulators

The third class of function regulators are the integrin to other molecule regulators. These compositions are designed to specifically interfere with molecules such as small molecule ligands or other proteins that interact with the integrin or integrin receptor. For example, an integrin to other molecule regulator might target a ligand for a particular integrin receptor, such as the alpha6beta4 receptor. The ligand for the alpha6beta4 receptor is laminin5 or the laminin gamma2 chain. Compositions that interact with laminin5 such that laminin5 or the laminin gamma2 chain interactions with alpha6beta4 are inhibited or reduced are specifically contemplated herein.

Likewise, there are other molecules, such as Shc molecules, that also interact with integrins, such as the alpha6beta4 receptor. Compositions that specifically interact with the Shc molecules such that they prevent the appropriate interactions between the Shc molecule and the alpha6beta4

receptor are disclosed.

(2) Types

Just as there are different general classes of molecules that can regulate (such as by inhibition) the function of the disclosed integrins and integrin receptors, so to there are many different types of molecules that perform that regulation. For example, any molecule that can perform the regulation of for example, the disclosed integrins, integrin receptors, or signaling pathways produced by the disclosed integrins and integrin receptors are contemplated. For example, antibodies or small molecules which inhibit the disclosed compositions are herein disclosed. Also disclosed are, for example, functional nucleic acids, such as ribozymes or antisense molecules that can inhibit the disclosed integrin function in a variety of ways. A non-limiting list of exemplary molecules is discussed herein.

(a) Antibodies

Antibodies can be used to regulate, for example, the function of the disclosed integrins and integrin receptors, molecules that interact with the disclosed integrin receptors, and molecules in the signaling pathways of the disclosed integrin receptors.

As used herein, the term "antibody" encompasses, but is not limited to, whole immunoglobulin (i.e., an intact antibody) of any class. Native antibodies are usually heterotetrameric glycoproteins, composed of two identical light (L) chains and two identical heavy (H) chains. Typically, each light chain is linked to a heavy chain by one covalent disulfide bond, 20 while the number of disulfide linkages varies between the heavy chains of different immunoglobulin isotypes. Each heavy and light chain also has regularly spaced intrachain disulfide bridges. Each heavy chain has at one end a variable domain (V(H)) followed by a number of constant domains. Each light chain has a variable domain at one end (V(L)) and a constant domain at its other end; the constant domain of the light chain is aligned with the first 25 constant domain of the heavy chain, and the light chain variable domain is aligned with the variable domain of the heavy chain. Particular amino acid residues are believed to form an interface between the light and heavy chain variable domains. The light chains of antibodies from any vertebrate species can be assigned to one of two clearly distinct types, called kappa (k) and lambda (1), based on the amino acid sequences of their constant domains. Depending on the amino 30 acid sequence of the constant domain of their heavy chains, immunoglobulins can be assigned to different classes. There are five major classes of human immunoglobulins: IgA, IgD, IgE, IgG and lgM, and several of these may be further divided into subclasses (isotypes), e.g., IgG-1, IgG-2, IgG-3, and IgG-4; IgA-1 and IgA-2. One skilled in the art would recognize the comparable classes for mouse. The heavy chain constant domains that correspond to the different classes of 35 immunoglobulins are called alpha, delta, epsilon, gamma, and mu, respectively.

Antibodies can be either polyclonal or monoclonal. Polyclonal antibodies, typically are derived from the serum of an animal that has been immunogenically challenged, and monoclonal antibodies are derived as discussed herein.

The term "variable" is used herein to describe certain portions of the variable domains that

differ in sequence among antibodies and are used in the binding and specificity of each particular antibody for its particular antigen. However, the variability is not usually evenly distributed through the variable domains of antibodies. It is typically concentrated in three segments called complementarity determining regions (CDRs) or hypervariable regions both in the light chain and the heavy chain variable domains. The more highly conserved portions of the variable domains are called the framework (FR). The variable domains of native heavy and light chains each comprise four FR regions, largely adopting a b-sheet configuration, connected by three CDRs, which form loops connecting, and in some cases forming part of, the b-sheet structure. The CDRs in each chain are held together in close proximity by the FR regions and, with the CDRs from the other chain, contribute to the formation of the antigen binding site of antibodies (see Kabat E. A. et al., "Sequences of Proteins of Immunological Interest," National Institutes of Health, Bethesda, Md. (1987)). The constant domains are not involved directly in binding an antibody to an antigen, but exhibit various effector functions, such as participation of the antibody in antibody-dependent cellular toxicity.

As used herein, the term "antibody or fragments thereof" encompasses chimeric antibodies and hybrid antibodies, with dual or multiple antigen or epitope specificities, and fragments, such as F(ab')2, Fab', Fab and the like, including hybrid fragments. Thus, fragments of the antibodies that retain the ability to bind their specific antigens are provided. For example, fragments of antibodies which maintain EphA2 binding activity are included within the meaning of the term "antibody or fragment thereof." Such antibodies and fragments can be made by techniques known in the art and can be screened for specificity and activity according to the methods set forth in the Examples and in general methods for producing antibodies and screening antibodies for specificity and activity (See Harlow and Lane. Antibodies, A Laboratory Manual. Cold Spring Harbor Publications, New York, (1988)).

Also included within the meaning of "antibody or fragments thereof" are conjugates of antibody fragments and antigen binding proteins (single chain antibodies) as described, for example, in U.S. Pat. No. 4,704,692, the contents of which are hereby incorporated by reference. Single chain divalent antibodies are also provided.

Optionally, the antibodies are generated in other species and "humanized" for administration in humans. Humanized forms of non-human (e.g., murine) antibodies are chimeric immunoglobulins, immunoglobulin chains or fragments thereof (such as Fv, Fab, Fab', F(ab')2, or

other antigen-binding subsequences of antibodies) which contain minimal sequence derived from non-human immunoglobulin. Humanized antibodies include human immunoglobulins (recipient antibody) in which residues from a complementary determining region (CDR) of the recipient are replaced by residues from a CDR of a non-human species (donor antibody) such as mouse, rat or rabbit having the desired specificity, affinity and capacity. In some instances, Fv framework residues of the human immunoglobulin are replaced by corresponding non-human residues. Humanized antibodies may also comprise residues that are found neither in the recipient antibody nor in the imported CDR or framework sequences. In general, the humanized antibody will comprise substantially all of at least one, and typically two, variable domains, in which all or substantially all of the CDR regions correspond to those of a non-human immunoglobulin and all or substantially all of the FR regions are those of a human immunoglobulin consensus sequence. The humanized antibody optimally also will comprise at least a portion of an immunoglobulin constant region (Fc), typically that of a human immunoglobulin (Jones et al., Nature, 321:522-525 (1986); Riechmann et al., Nature, 332:323-327 (1988); and Presta, Curr. Op. Struct. Biol., 2:593-596 (1992)).

Methods for humanizing non-human antibodies are well known in the art. Generally, a humanized antibody has one or more amino acid residues introduced into it from a source that is non-human. These non-human amino acid residues are often referred to as "import" residues, which are typically taken from an "import" variable domain. Humanization can be essentially performed following the method of Winter and co-workers (Jones et al., Nature, 321:522-525 (1986); Riechmann et al., Nature, 332:323-327 (1988); Verhoeyen et al., Science, 239:1534-1536 (1988)), by substituting rodent CDRs or CDR sequences for the corresponding sequences of a human antibody. Accordingly, such "humanized" antibodies are chimeric antibodies (U.S. Pat. No. 4,816,567), wherein substantially less than an intact human variable domain has been substituted by the corresponding sequence from a non-human species. In practice, humanized antibodies are typically human antibodies in which some CDR residues and possibly some FR residues are substituted by residues from analogous sites in rodent antibodies.

The choice of human variable domains, both light and heavy, to be used in making the humanized antibodies is very important in order to reduce antigenicity. According to the "best-fit" method, the sequence of the variable domain of a rodent antibody is screened against the entire library of known human variable domain sequences. The human sequence which is closest to that of the rodent is then accepted as the human framework (FR) for the humanized antibody (Sims et al., J. Immunol., 151:2296 (1993) and Chothia et al., J. Mol. Biol., 196:901 (1987)). Another method uses a particular framework derived from the consensus sequence of all human antibodies of a particular subgroup of light or heavy chains. The same framework may be used for several different humanized antibodies (Carter et al., Proc. Natl. Acad. Sci. USA, 89:4285 (1992);

Presta et al., J. Immunol., 151:2623 (1993)).

It is further important that antibodies be humanized with retention of high affinity for the antigen and other favorable biological properties. To achieve this goal, according to a preferred method, humanized antibodies are prepared by a process of analysis of the parental sequences and various conceptual humanized products using three dimensional models of the parental and humanized sequences. Three dimensional immunoglobulin models are commonly available and are familiar to those skilled in the art. Computer programs are available which illustrate and display probable three-dimensional conformational structures of selected candidate immunoglobulin sequences. Inspection of these displays permits analysis of the likely role of the residues in the functioning of the candidate immunoglobulin sequence, i.e., the analysis of residues that influence the ability of the candidate immunoglobulin to bind its antigen. In this way, FR residues can be selected and combined from the consensus and import sequence so that the desired antibody characteristic, such as increased affinity for the target antigen(s), is achieved. In general, the CDR residues are directly and most substantially involved in influencing antigen binding (see, WO 94/04679, published 3 March 1994).

Transgenic animals (e.g., mice) that are capable, upon immunization, of producing a full repertoire of human antibodies in the absence of endogenous immunoglobulin production can be employed. For example, it has been described that the homozygous deletion of the antibody heavy chain joining region (J(H)) gene in chimeric and germ-line mutant mice results in complete inhibition of endogenous antibody production. Transfer of the human germ-line immunoglobulin gene array in such germ-line mutant mice will result in the production of human antibodies upon antigen challenge (see, e.g., Jakobovits et al., Proc. Natl. Acad. Sci. USA, 90:2551-255 (1993); Jakobovits et al., Nature, 362:255-258 (1993); Bruggemann et al., Year in Immuno., 7:33 (1993)). Human antibodies can also be produced in phage display libraries (Hoogenboom et al., J. Mol. Biol., 227:381 (1991); Marks et al., J. Mol. Biol., 222:581 (1991)). The techniques of Cote et al. and Boerner et al. are also available for the preparation of human monoclonal antibodies (Cole et al., Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, p. 77 (1985); Boerner et al., J. Immunol., 147(1):86-95 (1991)).

The present invention further provides a hybridoma cell that produces the monoclonal antibody of the invention. The term "monoclonal antibody" as used herein refers to an antibody obtained from a substantially homogeneous population of antibodies, i.e., the individual antibodies comprising the population are identical except for possible naturally occurring mutations that may be present in minor amounts. The monoclonal antibodies herein specifically include "chimeric" antibodies in which a portion of the heavy and/or light chain is identical with or homologous to corresponding sequences in antibodies derived from a particular species or belonging to a

particular antibody class or subclass, while the remainder of the chain(s) is identical with or homologous to corresponding sequences in antibodies derived from another species or belonging to another antibody class or subclass, as well as fragments of such antibodies, so long as they exhibit the desired activity (See, U.S. Pat. No. 4,816,567 and Morrison et al., Proc. Natl. Acad. Sci. USA, 81:6851-6855 (1984)).

Monoclonal antibodies of the invention may be prepared using hybridoma methods, such as those described by Kohler and Milstein, Nature, 256:495 (1975) or Harlow and Lane. Antibodies, A Laboratory Manual. Cold Spring Harbor Publications, New York, (1988). In a hybridoma method, a mouse or other appropriate host animal, is typically immunized with an 10 immunizing agent to elicit lymphocytes that produce or are capable of producing antibodies that will specifically bind to the immunizing agent. Alternatively, the lymphocytes may be immunized in vitro. Preferably, the immunizing agent comprises EphA2. Traditionally, the generation of monoclonal antibodies has depended on the availability of purified protein or peptides for use as the immunogen. More recently DNA based immunizations have shown promise as a way to elicit 15 strong immune responses and generate monoclonal antibodies. In this approach, DNA-based immunization can be used, wherein DNA encoding a portion of EphA2 expressed as a fusion protein with human IgG1 is injected into the host animal according to methods known in the art (e.g., Kilpatrick KE, et al. Gene gun delivered DNA-based immunizations mediate rapid production of murine monoclonal antibodies to the Flt-3 receptor. Hybridoma. 1998 20 Dec;17(6):569-76; Kilpatrick KE et al. High-affinity monoclonal antibodies to PED/PEA-15 generated using 5 microg of DNA. Hybridoma. 2000 Aug;19(4):297-302, which are incorporated herein by referenced in full for the methods of antibody production).

An alternate approach to immunizations with either purified protein or DNA is to use antigen expressed in baculovirus. The advantages to this system include ease of generation, high levels of expression, and post-translational modifications that are highly similar to those seen in mammalian systems. Use of this system involves expressing domains of EphA2 antibody as fusion proteins. The antigen is produced by inserting a gene fragment in-frame between the signal sequence and the mature protein domain of the EphA2 antibody nucleotide sequence. This results in the display of the foreign proteins on the surface of the virion. This method allows immunization with whole virus, eliminating the need for purification of target antigens.

Generally, peripheral blood lymphocytes ("PBLs") are used in methods of producing monoclonal antibodies if cells of human origin are desired, or spleen cells or lymph node cells are used if non-human mammalian sources are desired. The lymphocytes are then fused with an immortalized cell line using a suitable fusing agent, such as polyethylene glycol, to form a hybridoma cell (Goding, "Monoclonal Antibodies: Principles and Practice" Academic Press,

(1986) pp. 59-103). Immortalized cell lines are usually transformed mammalian cells, including myeloma cells of rodent, bovine, equine, and human origin. Usually, rat or mouse myeloma cell lines are employed. The hybridoma cells may be cultured in a suitable culture medium that preferably contains one or more substances that inhibit the growth or survival of the unfused, 5 immortalized cells. For example, if the parental cells lack the enzyme hypoxanthine guanine phosphoribosyl transferase (HGPRT or HPRT), the culture medium for the hybridomas typically will include hypoxanthine, aminopterin, and thymidine ("HAT medium"), which substances prevent the growth of HGPRT-deficient cells. Preferred immortalized cell lines are those that fuse efficiently, support stable high level expression of antibody by the selected antibody-producing 10 cells, and are sensitive to a medium such as HAT medium. More preferred immortalized cell lines are murine myeloma lines, which can be obtained, for instance, from the Salk Institute Cell Distribution Center, San Diego, Calif. and the American Type Culture Collection, Rockville, Md. Human myeloma and mouse-human heteromyeloma cell lines also have been described for the production of human monoclonal antibodies (Kozbor, J. Immunol., 133:3001 (1984); Brodeur et 15 al., "Monoclonal Antibody Production Techniques and Applications" Marcel Dekker, Inc., New York, (1987) pp. 51-63).

The culture medium in which the hybridoma cells are cultured can then be assayed for the presence of monoclonal antibodies directed against EphA2. Preferably, the binding specificity of monoclonal antibodies produced by the hybridoma cells is determined by immunoprecipitation or by an in vitro binding assay, such as radioimmunoassay (RIA) or enzyme-linked immunoabsorbent assay (ELISA). Such techniques and assays are known in the art, and are described further in the Examples below or in Harlow and Lane "Antibodies, A Laboratory Manual" Cold Spring Harbor Publications, New York, (1988).

After the desired hybridoma cells are identified, the clones may be subcloned by limiting

25 dilution or FACS sorting procedures and grown by standard methods. Suitable culture media for
this purpose include, for example, Dulbecco's Modified Eagle's Medium and RPMI-1640 medium.

Alternatively, the hybridoma cells may be grown in vivo as ascites in a mammal.

The monoclonal antibodies secreted by the subclones may be isolated or purified from the culture medium or ascites fluid by conventional immunoglobulin purification procedures such as, for example, protein A-Sepharose, protein G, hydroxylapatite chromatography, gel electrophoresis, dialysis, or affinity chromatography.

The monoclonal antibodies may also be made by recombinant DNA methods, such as those described in U.S. Pat. No. 4,816,567. DNA encoding the monoclonal antibodies of the invention can be readily isolated and sequenced using conventional procedures (e.g., by using oligonucleotide probes that are capable of binding specifically to genes encoding the heavy and

light chains of murine antibodies). The hybridoma cells of the invention serve as a preferred source of such DNA. Once isolated, the DNA may be placed into expression vectors, which are then transfected into host cells such as simian COS cells, Chinese hamster ovary (CHO) cells, plasmacytoma cells, or myeloma cells that do not otherwise produce immunoglobulin protein, to obtain the synthesis of monoclonal antibodies in the recombinant host cells. The DNA also may be modified, for example, by substituting the coding sequence for human heavy and light chain constant domains in place of the homologous murine sequences (U.S. Pat. No. 4,816,567) or by covalently joining to the immunoglobulin coding sequence all or part of the coding sequence for a non-immunoglobulin polypeptide. Optionally, such a non-immunoglobulin polypeptide is substituted for the constant domains of an antibody of the invention or substituted for the variable domains of one antigen-combining site of an antibody of the invention to create a chimeric bivalent antibody comprising one antigen-combining site having specificity for EphA2 and another antigen-combining site having specificity for a different antigen.

In vitro methods are also suitable for preparing monovalent antibodies. Digestion of
antibodies to produce fragments thereof, particularly, Fab fragments, can be accomplished using
routine techniques known in the art. For instance, digestion can be performed using papain.
Examples of papain digestion are described in WO 94/29348 published Dec. 22, 1994, U.S. Pat.
No. 4,342,566, and Harlow and Lane, Antibodies, A Laboratory Manual, Cold Spring Harbor
Publications, New York, (1988). Papain digestion of antibodies typically produces two identical
antigen binding fragments, called Fab fragments, each with a single antigen binding site, and a
residual Fc fragment. Pepsin treatment yields a fragment, called the F(ab')2 fragment, that has
two antigen combining sites and is still capable of cross-linking antigen.

The Fab fragments produced in the antibody digestion also contain the constant domains of the light chain and the first constant domain of the heavy chain. Fab' fragments differ from Fab fragments by the addition of a few residues at the carboxy terminus of the heavy chain domain including one or more cysteines from the antibody hinge region. The F(ab')2 fragment is a bivalent fragment comprising two Fab' fragments linked by a disulfide bridge at the hinge region. Fab'-SH is the designation herein for Fab' in which the cysteine residue(s) of the constant domains bear a free thiol group. Antibody fragments originally were produced as pairs of Fab' fragments which have hinge cysteines between them. Other chemical couplings of antibody fragments are also known.

An isolated immunogenically specific paratope or fragment of the antibody is also provided. A specific immunogenic epitope of the antibody can be isolated from the whole antibody by chemical or mechanical disruption of the molecule. The purified fragments thus obtained are tested to determine their immunogenicity and specificity by the methods taught

herein. Immunoreactive paratopes of the antibody, optionally, are synthesized directly. An immunoreactive fragment is defined as an amino acid sequence of at least about two to five consecutive amino acids derived from the antibody amino acid sequence.

One method of producing proteins comprising the antibodies of the present invention is to 5 link two or more peptides or polypeptides together by protein chemistry techniques. For example, peptides or polypeptides can be chemically synthesized using currently available laboratory equipment using either Fmoc (9-fluorenylmethyloxycarbonyl) or Boc (tert -butyloxycarbonoyl) chemistry. (Applied Biosystems, Inc., Foster City, CA). One skilled in the art can readily appreciate that a peptide or polypeptide corresponding to the antibody of the present invention, for 10 example, can be synthesized by standard chemical reactions. For example, a peptide or polypeptide can be synthesized and not cleaved from its synthesis resin whereas the other fragment of an antibody can be synthesized and subsequently cleaved from the resin, thereby exposing a terminal group which is functionally blocked on the other fragment. By peptide condensation reactions, these two fragments can be covalently joined via a peptide bond at their carboxyl and 15 amino termini, respectively, to form an antibody, or fragment thereof. (Grant GA (1992) Synthetic Peptides: A User Guide. W.H. Freeman and Co., N.Y. (1992); Bodansky M and Trost B., Ed. (1993) Principles of Peptide Synthesis. Springer-Verlag Inc., NY. Alternatively, the peptide or polypeptide is independently synthesized in vivo as described above. Once isolated, these independent peptides or polypeptides may be linked to form an antibody or fragment thereof via 20 similar peptide condensation reactions.

For example, enzymatic ligation of cloned or synthetic peptide segments allow relatively short peptide fragments to be joined to produce larger peptide fragments, polypeptides or whole protein domains (Abrahmsen L et al., Biochemistry, 30:4151 (1991)). Alternatively, native chemical ligation of synthetic peptides can be utilized to synthetically construct large peptides or polypeptides from shorter peptide fragments. This method consists of a two step chemical reaction (Dawson et al. Synthesis of Proteins by Native Chemical Ligation. Science, 266:776-779 (1994)). The first step is the chemoselective reaction of an unprotected synthetic peptide-alpha-thioester with another unprotected peptide segment containing an amino-terminal Cys residue to give a thioester-linked intermediate as the initial covalent product. Without a change in the reaction conditions, this intermediate undergoes spontaneous, rapid intramolecular reaction to form a native peptide bond at the ligation site. Application of this native chemical ligation method to the total synthesis of a protein molecule is illustrated by the preparation of human interleukin 8 (IL-8) (Baggiolini M et al. (1992) FEBS Lett. 307:97-101; Clark-Lewis I et al., J.Biol.Chem., 269:16075 (1994); Clark-Lewis I et al., Biochemistry, 30:3128 (1991); Rajarathnam K et al., Biochemistry 33:6623-30 (1994)).

Alternatively, unprotected peptide segments are chemically linked where the bond formed between the peptide segments as a result of the chemical ligation is an unnatural (non-peptide) bond (Schnolzer, M et al. Science, 256:221 (1992)). This technique has been used to synthesize analogs of protein domains as well as large amounts of relatively pure proteins with full biological activity (deLisle Milton RC et al., Techniques in Protein Chemistry IV. Academic Press, New York, pp. 257-267 (1992)).

The invention also provides fragments of antibodies which have bioactivity. The polypeptide fragments of the present invention can be recombinant proteins obtained by cloning nucleic acids encoding the polypeptide in an expression system capable of producing the 10 polypeptide fragments thereof, such as an adenovirus or baculovirus expression system. For example, one can determine the active domain of an antibody from a specific hybridoma that can cause a biological effect associated with the interaction of the antibody with EphA2. For example, amino acids found to not contribute to either the activity or the binding specificity or affinity of the antibody can be deleted without a loss in the respective activity. For example, in various 15 embodiments, amino or carboxy-terminal amino acids are sequentially removed from either the native or the modified non-immunoglobulin molecule or the immunoglobulin molecule and the respective activity assayed in one of many available assays. In another example, a fragment of an antibody comprises a modified antibody wherein at least one amino acid has been substituted for the naturally occurring amino acid at a specific position, and a portion of either amino terminal or 20 carboxy terminal amino acids, or even an internal region of the antibody, has been replaced with a polypeptide fragment or other moiety, such as biotin, which can facilitate in the purification of the modified antibody. For example, a modified antibody can be fused to a maltose binding protein, through either peptide chemistry or cloning the respective nucleic acids encoding the two polypeptide fragments into an expression vector such that the expression of the coding region 25 results in a hybrid polypeptide. The hybrid polypeptide can be affinity purified by passing it over an amylose affinity solid support, and the modified antibody receptor can then be separated from the maltose binding region by cleaving the hybrid polypeptide with the specific protease factor Xa. (See, for example, New England Biolabs Product Catalog, 1996, pg. 164.). Similar purification procedures are available for isolating hybrid proteins from eukaryotic cells as well.

The fragments, whether attached to other sequences or not, include insertions, deletions, substitutions, or other selected modifications of particular regions or specific amino acids residues, provided the activity of the fragment is not significantly altered or impaired compared to the nonmodified antibody or antibody fragment. These modifications can provide for some additional property, such as to remove or add amino acids capable of disulfide bonding, to increase its biolongevity, to alter its secretory characteristics, etc. In any case, the fragment must possess a bioactive property, such as binding activity, regulation of binding at the binding domain, etc.

Functional or active regions of the antibody may be identified by mutagenesis of a specific region of the protein, followed by expression and testing of the expressed polypeptide. Such methods are readily apparent to a skilled practitioner in the art and can include site-specific mutagenesis of the nucleic acid encoding the antigen. (Zoller MJ et al. Nucl. Acids Res. 10:6487-500 (1982).

A variety of immunoassay formats may be used to select antibodies that selectively bind with a particular protein, variant, or fragment. For example, solid-phase ELISA immunoassays are routinely used to select antibodies selectively immunoreactive with a protein, protein variant, or fragment thereof. See Harlow and Lane. Antibodies, A Laboratory Manual. Cold Spring Harbor Publications, New York, (1988), for a description of immunoassay formats and conditions that could be used to determine selective binding. The binding affinity of a monoclonal antibody can, for example, be determined by the Scatchard analysis of Munson et al., Anal. Biochem., 107:220 (1980).

Also provided is an antibody reagent kit comprising containers of the monoclonal antibody or fragment thereof of the invention and one or more reagents for detecting binding of the antibody or fragment thereof to the EphA2 receptor molecule. The reagents can include, for example, fluorescent tags, enzymatic tags, or other tags. The reagents can also include secondary or tertiary antibodies or reagents for enzymatic reactions, wherein the enzymatic reactions produce a product that can be visualized.

(b) Functional Nucleic Acids

Functional nucleic acids can also be used to regulate the, for example, the function of the disclosed integrins, integrin receptors, molecules that interact with the disclosed integrin receptors, and molecules in the signaling pathways of the disclosed integrin receptors.

Functional nucleic acids are nucleic acid molecules that have a specific function, such as binding a target molecule or catalyzing a specific reaction. Functional nucleic acid molecules can be divided into the following categories, which are not meant to be limiting. For example, functional nucleic acids include antisense molecules, aptamers, ribozymes, triplex forming molecules, and external guide sequences. The functional nucleic acid molecules can act as affectors, inhibitors, modulators, and stimulators of a specific activity possessed by a target molecule, or the functional nucleic acid molecules can possess a de novo activity independent of any other molecules.

Functional nucleic acid molecules can interact with any macromolecule, such as DNA, RNA, polypeptides, or carbohydrate chains. Thus, functional nucleic acids can interact with the mRNA of beta4 integrin for example, or the genomic DNA of alpha6 integrin for example, or they can interact with the polypeptide laminin5, or the gamma2 subunit of laminin5. Often functional nucleic acids are designed to interact with other nucleic acids based on sequence homology

between the target molecule and the functional nucleic acid molecule. In other situations, the specific recognition between the functional nucleic acid molecule and the target molecule is not based on sequence homology between the functional nucleic acid molecule and the target molecule, but rather is based on the formation of tertiary structure that allows specific recognition 5 to take place. Antisense molecules are designed to interact with a target nucleic acid molecule through either canonical or non-canonical base pairing. The interaction of the antisense molecule and the target molecule is designed to promote the destruction of the target molecule through, for example, RNAseH mediated RNA-DNA hybrid degradation. Alternatively the antisense molecule is designed to interrupt a processing function that normally would take place on the target 10 molecule, such as transcription or replication. Antisense molecules can be designed based on the sequence of the target molecule. Numerous methods for optimization of antisense efficiency by finding the most accessible regions of the target molecule exist. Exemplary methods would be in vitro selection experiments and DNA modification studies using DMS and DEPC. It is preferred that antisense molecules bind the target molecule with a dissociation constant (k_d)less than 10-6. It 15 is more preferred that antisense molecules bind with a k_d less than 10-8. It is also more preferred that the antisense molecules bind the target molecule with a k_d less than 10^{-10} . It is also preferred that the antisense molecules bind the target molecule with a k_d less than 10⁻¹². A representative sample of methods and techniques which aid in the design and use of antisense molecules can be found in the following non-limiting list of United States patents: 5,135,917, 5,294,533, 5,627,158, 20 5,641,754, 5,691,317, 5,780,607, 5,786,138, 5,849,903, 5,856,103, 5,919,772, 5,955,590, 5,990,088, 5,994,320, 5,998,602, 6,005,095, 6,007,995, 6,013,522, 6,017,898, 6,018,042, 6,025,198, 6,033,910, 6,040,296, 6,046,004, 6,046,319, and 6,057,437.

Aptamers are molecules that interact with a target molecule, preferably in a specific way.

Typically aptamers are small nucleic acids ranging from 15-50 bases in length that fold into

defined secondary and tertiary structures, such as stem-loops or G-quartets. Aptamers can bind small molecules, such as ATP (United States patent 5,631,146) and theophiline (United States patent 5,580,737), as well as large molecules, such as reverse transcriptase (United States patent 5,786,462) and thrombin (United States patent 5,543,293). Aptamers can bind very tightly with k₀s from the target molecule of less than 10⁻¹² M. It is preferred that the aptamers bind the target molecule with a k₀ less than 10⁻⁶. It is more preferred that the aptamers bind the target molecule with a k₀ less than 10⁻¹⁶. It is also more preferred that the aptamers bind the target molecule with a k₀ less than 10⁻¹⁶. It is also preferred that the aptamers bind the target molecule with a k₀ less than 10⁻¹⁶. Aptamers can bind the target molecule with a very high degree of specificity. For example, aptamers have been isolated that have greater than a 10000 fold difference in binding affinities between the target molecule and another molecule that differ at only a single position on the molecule (United States patent 5,543,293). It is preferred that the aptamer have a k₀ with the target

molecule at least 10 fold lower than the k_d with a background binding molecule. It is more preferred that the aptamer have a k_d with the target molecule at least 100 fold lower than the k_d with a background binding molecule. It is more preferred that the aptamer have a k_d with the target molecule at least 1000 fold lower than the k_d with a background binding molecule. It is preferred that the aptamer have a k_d with the target molecule at least 10000 fold lower than the k_d with a background binding molecule. It is preferred when doing the comparison for a polypeptide for example, that the background molecule be a different polypeptide. For example, when determining the specificity of beta4 integrin aptamers, the background protein could be bovine serum albumin. Representative examples of how to make and use aptamers to bind a variety of different target molecules can be found in the following non-limiting list of United States patents: 5,476,766, 5,503,978, 5,631,146, 5,731,424, 5,780,228, 5,792,613, 5,795,721, 5,846,713, 5,858,660, 5,861,254, 5,864,026, 5,869,641, 5,958,691, 6,001,988, 6,011,020, 6,013,443, 6,020,130, 6,028,186, 6,030,776, and 6,051,698.

Ribozymes are nucleic acid molecules that are capable of catalyzing a chemical reaction, 15 either intramolecularly or intermolecularly. Ribozymes are thus catalytic nucleic acid. It is preferred that the ribozymes catalyze intermolecular reactions. There are a number of different types of ribozymes that catalyze nuclease or nucleic acid polymerase type reactions which are based on ribozymes found in natural systems, such as hammerhead ribozymes, (for example, but not limited to the following United States patents: 5,334,711, 5,436,330, 5,616,466, 5,633,133, 20 5,646,020, 5,652,094, 5,712,384, 5,770,715, 5,856,463, 5,861,288, 5,891,683, 5,891,684, 5,985,621, 5,989,908, 5,998,193, 5,998,203, WO 9858058 by Ludwig and Sproat, WO 9858057 by Ludwig and Sproat, and WO 9718312 by Ludwig and Sproat) hairpin ribozymes (for example, but not limited to the following United States patents: 5,631,115, 5,646,031, 5,683,902, 5,712,384, 5,856,188, 5,866,701, 5,869,339, and 6,022,962), and tetrahymena ribozymes (for example, but not 25 limited to the following United States patents: 5,595,873 and 5,652,107). There are also a number of ribozymes that are not found in natural systems, but which have been engineered to catalyze specific reactions de novo (for example, but not limited to the following United States patents: 5,580,967, 5,688,670, 5,807,718, and 5,910,408). Preferred ribozymes cleave RNA or DNA substrates, and more preferably cleave RNA substrates. Ribozymes typically cleave nucleic acid 30 substrates through recognition and binding of the target substrate with subsequent cleavage. This recognition is often based mostly on canonical or non-canonical base pair interactions. This property makes ribozymes particularly good candidates for target specific cleavage of nucleic acids because recognition of the target substrate is based on the target substrates sequence. Representative examples of how to make and use ribozymes to catalyze a variety of different 35 reactions can be found in the following non-limiting list of United States patents: 5,646,042, 5,693,535, 5,731,295, 5,811,300, 5,837,855, 5,869,253, 5,877,021, 5,877,022, 5,972,699,

5,972,704, 5,989,906, and 6,017,756.

Triplex forming functional nucleic acid molecules are molecules that can interact with either double-stranded or single-stranded nucleic acid. When triplex molecules interact with a target region, a structure called a triplex is formed, in which there are three strands of DNA forming a complex dependant on both Watson-Crick and Hoogsteen base-pairing. Triplex molecules are preferred because they can bind target regions with high affinity and specificity. It is preferred that the triplex forming molecules bind the target molecule with a k_d less than 10°. It is more preferred that the triplex forming molecules bind with a k_d less than 10°. It is also more preferred that the triplex forming molecules bind the target molecule with a k_d less than 10°. It is also preferred that the triplex forming molecules bind the target molecule with a k_d less than 10°. Representative examples of how to make and use triplex forming molecules to bind a variety of different target molecules can be found in the following non-limiting list of United States patents: 5,176,996, 5,645,985, 5,650,316, 5,683,874, 5,693,773, 5,834,185, 5,869,246, 5,874,566, and 5,962,426.

External guide sequences (EGSs) are molecules that bind a target nucleic acid molecule forming a complex, and this complex is recognized by RNase P, which cleaves the target molecule. EGSs can be designed to specifically target a RNA molecule of choice. RNAse P aids in processing transfer RNA (tRNA) within a cell. Bacterial RNAse P can be recruited to cleave virtually any RNA sequence by using an EGS that causes the target RNA:EGS complex to mimic the natural tRNA substrate. (WO 92/03566 by Yale, and Forster and Altman, Science 238:407-409 (1990)).

Similarly, eukaryotic EGS/RNAse P-directed cleavage of RNA can be utilized to cleave desired targets within eukaryotic cells. (Yuan et al., Proc. Natl. Acad. Sci. USA 89:8006-8010 (1992); WO 93/22434 by Yale; WO 95/24489 by Yale; Yuan and Altman, EMBO J 14:159-168 (1995), and Carrara et al., Proc. Natl. Acad. Sci. (USA) 92:2627-2631 (1995)). Representative examples of how to make and use EGS molecules to facilitate cleavage of a variety of different target molecules be found in the following non-limiting list of United States patents: 5,168,053, 5,624,824, 5,683,873, 5,728,521, 5,869,248, and 5,877,162

(c) Small molecules

30 Small molecules can also be used to regulate, for example, the function of the disclosed integrins, integrin receptors, molecules that interact with the disclosed integrin receptors, and molecules in the signaling pathways of the disclosed integrin receptors. Those of skill in the art understand how to generate small molecules of this type, and exemplary libraries and methods for isolating small molecule regulators are disclosed herein.

b) Compositions identified by screening with disclosed compositions / combinatorial chemistry

(1) Combinatorial chemistry

The disclosed compositions can be used as targets for any combinatorial technique to

identify molecules or macromolecular molecules that interact with the disclosed compositions,
such as beta4 integrin, alpha6 integrin, or the gamma2 subunit of laminin5, in a desired way. The
nucleic acids, peptides, and related molecules disclosed herein can be used as targets for the
combinatorial approaches. Also disclosed are the compositions that are identified through
combinatorial techniques or screening techniques in which the herein disclosed compositions, for
example set forth in SEQ ID NOS:1-19 or portions thereof, are used as the target or reagent in a
combinatorial or screening protocol.

It is understood that when using the disclosed compositions in combinatorial techniques or screening methods, molecules, such as macromolecular molecules, will be identified that have particular desired properties such as inhibition or stimulation or the target molecule's function.

The molecules identified and isolated when using the disclosed compositions, such as, beta4 integrin, alpa6 integrin, or the gamma2 subunit of laminin5, are also disclosed. Thus, the products produced using the combinatorial or screening approaches that involve the disclosed compositions, such as, beta4 integrin, alpha6 integrin, or the gamma2 subunit of laminin5, are also considered herein disclosed.

Combinatorial chemistry includes but is not limited to all methods for isolating small 20 molecules or macromolecules that are capable of binding either a small molecule or another macromolecule, typically in an iterative process. Proteins, oligonucleotides, and sugars are examples of macromolecules. For example, oligonucleotide molecules with a given function, catalytic or ligand-binding, can be isolated from a complex mixture of random oligonucleotides in 25 what has been referred to as "in vitro genetics" (Szostak, TIBS 19:89, 1992). One synthesizes a large pool of molecules bearing random and defined sequences and subjects that complex mixture, for example, approximately 1015 individual sequences in 100 µg of a 100 nucleotide RNA, to some selection and enrichment process. Through repeated cycles of affinity chromatography and PCR amplification of the molecules bound to the ligand on the solid support, Ellington and Szostak 30 (1990) estimated that 1 in 1010 RNA molecules folded in such a way as to bind a small molecule dyes. DNA molecules with such ligand-binding behavior have been isolated as well (Ellington and Szostak, 1992; Bock et al. 1992). Techniques aimed at similar goals exist for small organic molecules, proteins, antibodies and other macromolecules known to those of skill in the art. Screening sets of molecules for a desired activity whether based on small organic libraries, 35 oligonucleotides, or antibodies is broadly referred to as combinatorial chemistry. Combinatorial techniques are particularly suited for defining binding interactions between molecules and for

isolating molecules that have a specific binding activity, often called aptamers when the macromolecules are nucleic acids.

There are a number of methods for isolating proteins which either have de novo activity or a modified activity. For example, phage display libraries have been used to isolate numerous 5 peptides that interact with a specific target. (See for example, United States Patent No. 6,031,071; 5,824,520; 5,596,079; and 5,565,332 which are herein incorporated by reference at least for their material related to phage display and methods relate to combinatorial chemistry)

A preferred method for isolating proteins that have a given function is described by Roberts and Szostak (Roberts R.W. and Szostak J.W. Proc. Natl. Acad. Sci. USA, 10 94(23)12997-302 (1997). This combinatorial chemistry method couples the functional power of proteins and the genetic power of nucleic acids. An RNA molecule is generated in which a puromycin molecule is covalently attached to the 3'-end of the RNA molecule. An in vitro translation of this modified RNA molecule causes the correct protein, encoded by the RNA to be translated. In addition, because of the attachment of the puromycin, a peptdyl acceptor which 15 cannot be extended, the growing peptide chain is attached to the puromycin which is attached to the RNA. Thus, the protein molecule is attached to the genetic material that encodes it. Normal in vitro selection procedures can now be done to isolate functional peptides. Once the selection procedure for peptide function is complete traditional nucleic acid manipulation procedures are performed to amplify the nucleic acid that codes for the selected functional peptides. After 20 amplification of the genetic material, new RNA is transcribed with puromycin at the 3'-end, new peptide is translated and another functional round of selection is performed. Thus, protein selection can be performed in an iterative manner just like nucleic acid selection techniques. The peptide which is translated is controlled by the sequence of the RNA attached to the puromycin. This sequence can be anything from a random sequence engineered for optimum translation (i.e. 25 no stop codons etc.) or it can be a degenerate sequence of a known RNA molecule to look for improved or altered function of a known peptide. The conditions for nucleic acid amplification and in vitro translation are well known to those of ordinary skill in the art and are preferably performed as in Roberts and Szostak (Roberts R.W. and Szostak J.W. Proc. Natl. Acad. Sci. USA, 94(23)12997-302 (1997)).

Another preferred method for combinatorial methods designed to isolate peptides is described in Cohen et al. (Cohen B.A., et al., Proc. Natl. Acad. Sci. USA 95(24):14272-7 (1998)). This method utilizes and modifies two-hybrid technology. Yeast two-hybrid systems are useful for the detection and analysis of protein:protein interactions. The two-hybrid system, initially described in the yeast Saccharomyces cerevisiae, is a powerful molecular genetic technique for identifying new regulatory molecules, specific to the protein of interest (Fields and

Song, Nature 340:245-6 (1989)). Cohen et al., modified this technology so that novel interactions between synthetic or engineered peptide sequences could be identified which bind a molecule of choice. The benefit of this type of technology is that the selection is done in an intracellular environment. The method utilizes a library of peptide molecules that attached to an acidic activation domain. A peptide of choice, for example a portion of beta4 or alpha6 or gamma2 or laminin5 is attached to a DNA binding domain of a transcriptional activation protein, such as Gal 4. By performing the Two-hybrid technique on this type of system, molecules that bind the portion of beta4 or alpha6 or gamma2 or laminin5 can be identified.

There are molecules that can act like antibodies, in that they can having varying binding specificities, that are based on a fibronectin motif. The fibronectin type III domain (FN3) is a small autonomous folding unit. This FN3 domain can be found in numeorus proteins that bind ligand, such as animal proteins. The beta-sandwich structure of FN3 closely resembles that of immunoglobulin domains. FN3 mutants can be isolated using combinatorial approaches disclosed herein, for example phage display, that bind desired targets. Typically the libraries of FN3 molecules have been randomized in the two surface loops. Thus, FN3 can be used at least as a scaffold for engineering novel binding proteins. (Koide A, Bailey CW, Huang X, Koide S., "The fibronectin type III domain as a scaffold for novel binding proteins." J Mol Biol 1998: 284,1141-1151 which is herein incorporated by reference at least for material related to the fibronectin based novel binding proteins).

Using methodology well known to those of skill in the art, in combination with various combinatorial libraries, one can isolate and characterize those small molecules or macromolecules, which bind to or interact with the desired target. The relative binding affinity of these compounds can be compared and optimum compounds identified using competitive binding studies, which are well known to those of skill in the art.

Techniques for making combinatorial libraries and screening combinatorial libraries to isolate molecules which bind a desired target are well known to those of skill in the art.

Representative techniques and methods can be found in but are not limited to United States patents 5,084,824, 5,288,514, 5,449,754, 5,506,337, 5,539,083, 5,545,568, 5,556,762, 5,565,324, 5,565,332, 5,573,905, 5,618,825, 5,619,680, 5,627,210, 5,646,285, 5,663,046, 5,670,326, 30 5,677,195, 5,683,899, 5,688,696, 5,688,997, 5,698,685, 5,712,146, 5,721,099, 5,723,598, 5,741,713, 5,792,431, 5,807,683, 5,807,754, 5,821,130, 5,831,014, 5,834,195, 5,834,318, 5,834,588, 5,840,500, 5,847,150, 5,856,107, 5,856,496, 5,859,190, 5,864,010, 5,874,443, 5,877,214, 5,880,972, 5,886,126, 5,886,127, 5,891,737, 5,916,899, 5,919,955, 5,925,527, 5,939,268, 5,942,387, 5,945,070, 5,948,696, 5,958,702, 5,958,792, 5,962,337, 5,965,719, 35 5,972,719, 5,976,894, 5,980,704, 5,985,356, 5,999,086, 6,001,579, 6,004,617, 6,008,321,

6,017,768, 6,025,371, 6,030,917, 6,040,193, 6,045,671, 6,045,755, 6,060,596, and 6,061,636.

Combinatorial libraries can be made from a wide array of molecules using a number of different synthetic techniques. For example, libraries containing fused 2,4-pyrimidinediones (United States patent 6,025,371) dihydrobenzopyrans (United States Patent 6,017,768 and 5 5,821,130), amide alcohols (United States Patent 5,976,894), hydroxy-amino acid amides (United States Patent 5,972,719) carbohydrates (United States patent 5,965,719), 1,4-benzodiazepin-2,5diones (United States patent 5,962,337), cyclics (United States patent 5,958,792), biaryl amino acid amides (United States patent 5,948,696), thiophenes (United States patent 5,942,387), tricyclic Tetrahydroquinolines (United States patent 5,925,527), benzofurans (United States patent 10 5,919,955), isoquinolines (United States patent 5,916,899), hydantoin and thiohydantoin (United States patent 5,859,190), indoles (United States patent 5,856,496), imidazol-pyrido-indole and imidazol-pyrido-benzothiophenes (United States patent 5,856,107) substituted 2-methylene-2, 3dihydrothiazoles (United States patent 5,847,150), quinolines (United States patent 5,840,500), PNA (United States patent 5,831,014), containing tags (United States patent 5,721,099), 15 polyketides (United States patent 5,712,146), morpholino-subunits (United States patent 5,698,685 and 5,506,337), sulfamides (United States patent 5,618,825), and benzodiazepines (United States patent 5,288,514).

Screening molecules similar to alpha6 for inhibition of alpha6beta4 formation is a method of identifying and isolating desired compounds that can inhibit the formation of A6B4 receptor. 20 For example, the disclosed compositions, such as alpha6 integrin or beta4 integrin can be used as targets in a selection scheme disclosed herein, and then the counter part integrin could be used as a competitive inhibitor to isolate the desired molecules. For example, a library of molecules could be incubated with beta4 integrin, which is bound to a solid support. The solid support can be washed to remove the unbound molecules and then the solid support can be incubated with, for 25 example, alpha6 integrin at a concentration that will saturate all beta4 binding sites. The molecules which are collected in the flowthrough after washing the solid support will be enriched for molecules that interact with beta4 integrin in a way that is competitive to the alpha6-beta4 interaction. Likewise, the solid support, bound with a target integrin, or more preferably a target integrin receptor, such as alpha6beta4 receptor, could also be washed, with for example, laminin5 30 or the gamma2 subunit of laminin5 at a concentration that will saturate all of the gamma2 binding sites on the beta4 integrin. Collection of the wash under these conditions will yield a population of molecules enriched for molecules that competitively interact with beta4 integrin at the beta4gamma2 site. Another example, is the following: bind target to solid support on microtiter plate. Incubate with ligand in presence of gridded subset of library members (or single compounds), 35 wash, identify competitor by reduction of ligand binding. It is understood that the exemplary discussions of alpha6beta4 and/or beta4 are equally applicable to alpha6, as well as other

alpha6betax receptors, such alpha6beta1.

Also disclosed are methods of isolating molecules that bind with a target molecule selected from the group consisting, B4 integrin, alpha6 integrin, and the gamma2 subunit of laminin5 comprising 1) contacting a library of molecules with the target molecule and 2) collecting molecules that bind the target molecule producing an enriched population of molecules.

Disclosed are methods, further comprising the step of repeating steps 1 and 2 with the enriched population of molecules, and/or wherein the library comprises a small molecule, peptide, peptide mimetic, or oligonucleotide.

As used herein combinatorial methods and libraries included traditional screening methods and libraries as well as methods and libraries used in iterative processes.

(2) Computer assisted design

The disclosed compositions can be used as targets for any molecular modeling technique to identify either the structure of the disclosed compositions or to identify potential or actual molecules, such as small molecules, which interact in a desired way with the disclosed compositions. The nucleic acids, peptides, and related molecules disclosed herein can be used as targets in any molecular modeling program or approach.

It is understood that when using the disclosed compositions in modeling techniques, molecules, such as macromolecular molecules, will be identified that have particular desired properties such as inhibition or stimulation or the target molecule's function. The molecules identified and isolated when using the disclosed compositions, such as, beta4, are also disclosed. Thus, the products produced using the molecular modeling approaches that involve the disclosed compositions, such as, beta4 integrin, alpha6 integrin, or laminin5, are also considered herein disclosed.

Thus, one way to isolate molecules that bind a molecule of choice is through rational

design. This is achieved through structural information and computer modeling. Computer
modeling technology allows visualization of the three-dimensional atomic structure of a selected
molecule and the rational design of new compounds that will interact with the molecule. The
three-dimensional construct typically depends on data from x-ray crystallographic analyses or
NMR imaging of the selected molecule. The molecular dynamics require force field data. The
computer graphics systems enable prediction of how a new compound will link to the target
molecule and allow experimental manipulation of the structures of the compound and target
molecule to perfect binding specificity. Prediction of what the molecule-compound interaction
will be when small changes are made in one or both requires molecular mechanics software and
computationally intensive computers, usually coupled with user-friendly, menu-driven interfaces

between the molecular design program and the user.

Examples of molecular modeling systems are the CHARMm and QUANTA programs,
Polygen Corporation, Waltham, MA. CHARMm performs the energy minimization and molecular
dynamics functions. QUANTA performs the construction, graphic modeling and analysis of
molecular structure. QUANTA allows interactive construction, modification, visualization, and
analysis of the behavior of molecules with each other.

A number of articles review computer modeling of drugs interactive with specific proteins, such as Rotivinen, et al., 1988 Acta Pharmaceutica Fennica 97, 159-166; Ripka, New Scientist 54-57 (June 16, 1988); McKinaly and Rossmann, 1989 Annu. Rev. Pharmacol. Toxiciol. 29, 111-10 122; Perry and Davies, OSAR: Quantitative Structure-Activity Relationships in Drug Design pp. 189-193 (Alan R. Liss, Inc. 1989); Lewis and Dean, 1989 Proc. R. Soc. Lond. 236, 125-140 and 141-162; and, with respect to a model enzyme for nucleic acid components, Askew, et al., 1989 J. Am. Chem. Soc. 111, 1082-1090. Other computer programs that screen and graphically depict chemicals are available from companies such as BioDesign, Inc., Pasadena, CA., Allelix, Inc, Mississauga, Ontario, Canada, and Hypercube, Inc., Cambridge, Ontario. Although these are primarily designed for application to drugs specific to particular proteins, they can be adapted to design of molecules specifically interacting with specific regions of DNA or RNA, once that region is identified.

Although described above with reference to design and generation of compounds which could alter binding, one could also screen libraries of known compounds, including natural products or synthetic chemicals, and biologically active materials, including proteins, for compounds which alter substrate binding or enzymatic activity.

c) Nucleic acids

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There are a variety of molecules disclosed herein that are nucleic acid based, including for example the nucleic acids that encode, for example beta4 and the gamma2 subunit of laminin5, as well as various functional nucleic acids. The disclosed nucleic acids are made up of for example, nucleotides, nucleotide analogs, or nucleotide substitutes. Non-limiting examples of these and other molecules are discussed herein. It is understood that for example, when a vector is expressed in a cell, that the expressed mRNA will typically be made up of A, C, G, and U. Likewise, it is understood that if, for example, an antisense molecule is introduced into a cell or cell environment through for example exogenous delivery, it is advantageous that the antisense molecule be made up of nucleotide analogs that reduce the degradation of the antisense molecule in the cellular environment.

(1) Nucleotides and related molecules

A nucleotide is a molecule that contains a base moiety, a sugar moiety and a phosphate

moiety. Nucleotides can be linked together through their phosphate moieties and sugar moieties creating an internucleoside linkage. The base moiety of a nucleotide can be adenin-9-yl (A), cytosin-1-yl (C), guanin-9-yl (G), uracil-1-yl (U), and thymin-1-yl (T). The sugar moiety of a nucleotide is a ribose or a deoxyribose. The phosphate moiety of a nucleotide is pentavalent phosphate. An non-limiting example of a nucleotide would be 3'-AMP (3'-adenosine monophosphate) or 5'-GMP (5'-guanosine monophosphate).

A nucleotide analog is a nucleotide which contains some type of modification to either the base, sugar, or phosphate moieties. Modifications to the base moiety would include natural and synthetic modifications of A, C, G, and T/U as well as different purine or pyrimidine bases, such 10 as uracil-5-yl (.psi.), hypoxanthin-9-yl (I), and 2-aminoadenin-9-yl. A modified base includes but is not limited to 5-methylcytosine (5-me-C), 5-hydroxymethyl cytosine, xanthine, hypoxanthine, 2-aminoadenine, 6-methyl and other alkyl derivatives of adenine and guanine, 2-propyl and other alkyl derivatives of adenine and guanine, 2-thiouracil, 2-thiothymine and 2-thiocytosine, 5halouracil and cytosine, 5-propynyl uracil and cytosine, 6-azo uracil, cytosine and thymine, 5-15 uracil (pseudouracil), 4-thiouracil, 8-halo, 8-amino, 8-thiol, 8-thioalkyl, 8-hydroxyl and other 8substituted adenines and guanines, 5-halo particularly 5-bromo, 5-trifluoromethyl and other 5substituted uracils and cytosines, 7-methylguanine and 7-methyladenine, 8-azaguanine and 8azaadenine, 7-deazaguanine and 7-deazaadenine and 3-deazaguanine and 3-deazaadenine. Additional base modifications can be found for example in U.S. Pat. No. 3,687,808, Englisch et 20 al., Angewandte Chemie, International Edition, 1991, 30, 613, and Sanghvi, Y. S., Chapter 15, Antisense Research and Applications, pages 289-302, Crooke, S. T. and Lebleu, B. ed., CRC Press, 1993. Certain nucleotide analogs, such as 5-substituted pyrimidines, 6-azapyrimidines and N-2, N-6 and O-6 substituted purines, including 2-aminopropyladenine, 5-propynyluracil and 5propynylcytosine. 5-methylcytosine can increase the stability of duplex formation. Base 25 modifications can be combined, for example, with a sugar modification, such as 2'-Omethoxyethyl, to achieve unique properties such as increased duplex stability. There are numerous United States patents such as 4,845,205; 5,130,302; 5,134,066; 5,175,273; 5,367,066; 5,432,272; 5,457,187; 5,459,255; 5,484,908; 5,502,177; 5,525,711; 5,552,540; 5,587,469; 5,594,121, 5,596,091; 5,614,617; and 5,681,941, which detail and describe a range of base 30 modifications. Each of these patents is herein incorporated by reference.

Nucleotide analogs can also include modifications of the sugar moiety. Modifications to the sugar moiety would include natural modifications of the ribose and deoxy ribose as well as synthetic modifications. Sugar modifications include but are not limited to the following modifications at the 2' position: OH-; F-; O-, S-, or N-alkyl; O-, S-, or N-alkenyl; O-, S- or N-alkynyl; or O-alkyl-O-alkyl, wherein the alkyl, alkenyl and alkynyl may be substituted or unsubstituted C₁ to C₁₀, alkyl or C₂ to C₁₀ alkenyl and alkynyl. 2' sugar modifications also include

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but are not limited to $-O[(CH_2)_n O]_m CH_3$, $-O(CH_2)_n OCH_3$, $-O(CH_2)_n NH_2$, $-O(CH_2)_n CH_3$, $-O(CH_2)_n OCH_3$, and $-O(CH_2)_n ON[(CH_2)_n CH_3)]_2$, where n and m are from 1 to about 10.

Other modifications at the 2' position include but are not limited to: C1 to C10 lower alkyl, substituted lower alkyl, alkaryl, aralkyl, O-alkaryl or O-aralkyl, SH, SCH3, OCN, Cl, Br, CN, CF3, 5 OCF₃, SOCH₃, SO₂ CH₃, ONO₂, NO₂, N₃, NH₂, heterocycloalkyl, heterocycloalkaryl, aminoalkylamino, polyalkylamino, substituted silyl, an RNA cleaving group, a reporter group, an intercalator, a group for improving the pharmacokinetic properties of an oligonucleotide, or a group for improving the pharmacodynamic properties of an oligonucleotide, and other substituents having similar properties. Similar modifications may also be made at other positions on the sugar, 10 particularly the 3' position of the sugar on the 3' terminal nucleotide or in 2'-5' linked oligonucleotides and the 5' position of 5' terminal nucleotide. Modified sugars would also include those that contain modifications at the bridging ring oxygen, such as CH, and S. Nucleotide sugar analogs may also have sugar mimetics such as cyclobutyl moieties in place of the pentofuranosyl sugar. There are numerous United States patents that teach the preparation of such modified sugar 15 structures such as 4,981,957; 5,118,800; 5,319,080; 5,359,044; 5,393,878; 5,446,137; 5,466,786; 5,514,785; 5,519,134; 5,567,811; 5,576,427; 5,591,722; 5,597,909; 5,610,300; 5,627,053; 5,639,873; 5,646,265; 5,658,873; 5,670,633; and 5,700,920, each of which is herein incorporated by reference in its entirety.

Nucleotide analogs can also be modified at the phosphate moiety. Modified phosphate moieties include but are not limited to those that can be modified so that the linkage between two nucleotides contains a phosphorothioate, chiral phosphorothioate, phosphorodithioate, phosphotriester, aminoalkylphosphotriester, methyl and other alkyl phosphonates including 3'-alkylene phosphonate and chiral phosphonates, phosphinates, phosphoramidates including 3'-amino phosphoramidate and aminoalkylphosphoramidates, thionophosphoramidates, thionoalkylphosphonates, thionoalkylphosphoramidates, thionophosphates. It is understood that these phosphate or modified phosphate linkage between two nucleotides can be through a 3'-5' linkage or a 2'-5' linkage, and the linkage can contain inverted polarity such as 3'-5' to 5'-3' or 2'-5' to 5'-2'. Various salts, mixed salts and free acid forms are also included. Numerous United States patents teach how to make and use nucleotides containing modified phosphates and include but are not limited to, 3,687,808; 4,469,863; 4,476,301; 5,023,243; 5,177,196; 5,188,897; 5,264,423; 5,276,019; 5,278,302; 5,286,717; 5,321,131; 5,399,676; 5,405,939; 5,453,496; 5,455,233; 5,466,677; 5,476,925; 5,519,126; 5,536,821; 5,541,306; 5,550,111; 5,563,253; 5,571,799; 5,587,361; and 5,625,050, each of which is herein incorporated by reference.

It is understood that nucleotide analogs need only contain a single modification, but may also contain multiple modifications within one of the moieties or between different moieties.

Nucleotide substitutes are molecules having similar functional properties to nucleotides, but which do not contain a phosphate moiety, such as peptide nucleic acid (PNA). Nucleotide substitutes are molecules that will recognize nucleic acids in a Watson-Crick or Hoogsteen manner, but which are linked together through a moiety other than a phosphate moiety.

Nucleotide substitutes are able to conform to a double helix type structure when interacting with the appropriate target nucleic acid.

Nucleotide substitutes are nucleotides or nucleotide analogs that have had the phosphate moiety and/or sugar moieties replaced. Nucleotide substitutes do not contain a standard phosphorus atom. Substitutes for the phosphate can be for example, short chain alkyl or 10 cycloalkyl internucleoside linkages, mixed heteroatom and alkyl or cycloalkyl internucleoside linkages, or one or more short chain heteroatomic or heterocyclic internucleoside linkages. These include those having morpholino linkages (formed in part from the sugar portion of a nucleoside); siloxane backbones; sulfide, sulfoxide and sulfone backbones; formacetyl and thioformacetyl backbones; methylene formacetyl and thioformacetyl backbones; alkene containing backbones; 15 sulfamate backbones; methyleneimino and methylenehydrazino backbones; sulfonate and sulfonamide backbones; amide backbones; and others having mixed N, O, S and CH2 component parts. Numerous United States patents disclose how to make and use these types of phosphate replacements and include but are not limited to 5,034,506; 5,166,315; 5,185,444; 5,214,134; 5.216.141; 5.235,033; 5,264,562; 5,264,564; 5,405,938; 5,434,257; 5,466,677; 5,470,967; 20 5,489,677; 5,541,307; 5,561,225; 5,596,086; 5,602,240; 5,610,289; 5,602,240; 5,608,046; 5,610,289; 5,618,704; 5,623,070; 5,663,312; 5,633,360; 5,677,437; and 5,677,439, each of which is herein incorporated by reference.

It is also understood in a nucleotide substitute that both the sugar and the phosphate moieties of the nucleotide can be replaced, by for example an amide type linkage

25 (aminoethylglycine) (PNA). United States patents 5,539,082; 5,714,331; and 5,719,262 teach how to make and use PNA molecules, each of which is herein incorporated by reference. (See also Nielsen et al., Science, 1991, 254, 1497-1500).

It is also possible to link other types of molecules (conjugates) to nucleotides or nucleotide analogs to enhance for example, cellular uptake. Conjugates can be chemically linked to the nucleotide or nucleotide analogs. Such conjugates include but are not limited to lipid moieties such as a cholesterol moiety (Letsinger et al., Proc. Natl. Acad. Sci. USA, 1989,

86, 6553-6556), cholic acid (Manoharan et al., Bioorg. Med. Chem. Let., 1994, 4, 1053-1060), a thioether, e.g., hexyl-S-tritylthiol (Manoharan et al., Ann. N.Y. Acad. Sci., 1992, 660, 306-309; Manoharan et al., Bioorg. Med. Chem. Let., 1993, 3, 2765-2770), a thiocholesterol (Oberhauser et al., Nucl. Acids Res., 1992, 20, 533-538), an aliphatic chain, e.g., dodecandiol or

undecyl residues (Saison-Behmoaras et al., EMBO J., 1991, 10, 1111-1118; Kabanov et al., FEBS Lett., 1990, 259, 327-330; Svinarchuk et al., Biochimie, 1993, 75, 49-54), a phospholipid, e.g., dihexadecyl-rac-glycerol or triethylammonium 1,2-di-O-hexadecyl-rac-glycero-3-H-phosphonate (Manoharan et al., Tetrahedron Lett., 1995, 36, 3651-3654; Shea et al., Nucl. Acids Res., 1990, 5 18, 3777-3783), a polyamine or a polyethylene glycol chain (Manoharan et al., Nucleosides & Nucleotides, 1995, 14, 969-973), or adamantane acetic acid (Manoharan et al., Tetrahedron Lett., 1995, 36, 3651-3654), a palmityl moiety (Mishra et al., Biochim. Biophys. Acta, 1995, 1264, 229-237), or an octadecylamine or hexylamino-carbonyl-oxycholesterol moiety (Crooke et al., J. Pharmacol. Exp. Ther., 1996, 277, 923-937. Numerous United States patents teach the 10 preparation of such conjugates and include, but are not limited to U.S. Pat. Nos. 4,828,979; 4,948,882; 5,218,105; 5,525,465; 5,541,313; 5,545,730; 5,552,538; 5,578,717, 5,580,731; 5.580,731; 5.591,584; 5.109,124; 5.118,802; 5.138,045; 5,414,077; 5,486,603; 5,512,439; 5.578,718; 5.608,046; 4.587,044; 4,605,735; 4,667,025; 4,762,779; 4,789,737; 4,824,941; 4,835,263; 4,876,335; 4,904,582; 4,958,013; 5,082,830; 5,112,963; 5,214,136; 5,082,830; 15 5,112,963; 5,214,136; 5,245,022; 5,254,469; 5,258,506; 5,262,536; 5,272,250; 5,292,873; 5,317,098; 5,371,241, 5,391,723; 5,416,203, 5,451,463; 5,510,475; 5,512,667; 5,514,785; 5,565,552; 5,567,810; 5,574,142; 5,585,481; 5,587,371; 5,595,726; 5,597,696; 5,599,923; 5,599,928 and 5,688,941, each of which is herein incorporated by reference.

It is understood that an oligonucleotide can be made from any combination of nucleotides, nucleotide analogs, or nucleotide substitutes disclosed herein or related molecules not specifically recited herein.

A Watson-Crick interaction is at least one interaction with the Watson-Crick face of a nucleotide, nucleotide analog, or nucleotide substitute. The Watson-Crick face of a nucleotide, nucleotide analog, or nucleotide substitute includes the C2, N1, and C6 positions of a purine based nucleotide, nucleotide analog, or nucleotide substitute and the C2, N3, C4 positions of a pyrimidine based nucleotide, nucleotide analog, or nucleotide substitute.

A Hoogsteen interaction is the interaction that takes place on the Hoogsteen face of a nucleotide or nucleotide analog, which is exposed in the major groove of duplex DNA. The Hoogsteen face includes the N7 position and reactive groups (NH2 or O) at the C6 position of purine nucleotides.

(2) Sequences

There are a variety of sequences related to the beta4 integrin or the laminin5-gamma2 gene or the alpha6 integrin gene having the following Genbank Accession Numbers 6453379, 4557674, and AH006634, respectively. These sequences and others are herein incorporated by reference in their entireties as well as for individual subsequences contained therein.

One particular sequence set forth in SEQ ID NO:1 (beta4 integrin) and having Genbank accession number 6453379 is used herein, at various points, as an example, to exemplify the disclosed compositions and methods (or when another particular sequence is used as an example). It is understood that the description related to this sequence is applicable to any sequence related to beta4 integrin or any of the other molecules disclosed herein, such as alpha6 integrin, or the subunits of laminin5, such as gamma2, unless specifically indicated otherwise. Those of skill in the art understand how to resolve sequence discrepancies and differences and to adjust the compositions and methods relating to a particular sequence to other related sequences (i.e. sequences of beta4 integrin). Primers and/or probes can be designed for any beta4 integrin

(3) Primers and probes

Disclosed are compositions including primers and probes, which are capable of interacting with the for example, the alpha6 gene or mRNA, beta4 gene or mRNA, or gamma2 subunit of the laminin5 ligand gene as disclosed herein or mRNA as wells as primers or probes for any of the 15 sequences or fragments of the sequences, set forth in SEQ ID NOs:1,3,5,7,9,11, and 13. In certain embodiments the primers are used to support DNA amplification reactions. Typically the primers will be capable of being extended in a sequence specific manner. Extension of a primer in a sequence specific manner includes any methods wherein the sequence and/or composition of the nucleic acid molecule to which the primer is hybridized or otherwise associated directs or 20 influences the composition or sequence of the product produced by the extension of the primer. Extension of the primer in a sequence specific manner therefore includes, but is not limited to, PCR, DNA sequencing, DNA extension, DNA polymerization, RNA transcription, or reverse transcription. Techniques and conditions that amplify the primer in a sequence specific manner are preferred. In certain embodiments the primers are used for the DNA amplification reactions, 25 such as PCR or direct sequencing. It is understood that in certain embodiments the primers can also be extended using non-enzymatic techniques, where for example, the nucleotides or oligonucleotides used to extend the primer are modified such that they will chemically react to extend the primer in a sequence specific manner. Typically the disclosed primers hybridize with the beta4 gene, alpha6 gene, or laminin5-gamma2 subunit gene, for example, or region of the 30 beta4 gene, alpha6 gene, or laminin5-gamma2 subunit gene, for example, or they hybridize with the complement of the beta4 gene, alpha6 gene, or laminin5-gamma2 subunit gene, for example, or complement of a region of the beta4 gene, alpha6 gene, or laminin5-gamma2 subunit gene, or any of the sequences or fragments of the sequences, set forth in SEQ ID NOs:1,3,5,7,9,11, and 13, for example.

The size of the primers or probes for interaction with the beta4 gene, alpha6 gene, or laminin5-gamma2 subunit gene, for example, in certain embodiments can be any size that supports

the desired enzymatic manipulation of the primer, such as DNA amplification or the simple hybridization of the probe or primer. A typical primer or probe for beta4 gene, alpha6 gene, or laminin5-gamma2 subunit gene, for example, or primer or probe for any of the sequences or fragments of the sequences, set forth in SEQ ID NOs:1,3,5,7,9,11, and 13 would be at least about 5 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 125, 150, 175, 200, 225, 250, 275, 300, 325, 350, 375, 400, 425, 450, 475, 500, 550, 600, 650, 700, 750, 800, 850, 900, 950, 1000, 1250, 1500, 1750, 2000, 2250, 2500, 2750, 3000, 3500, or 4000 nucleotides long.

In other embodiments a primer or probe for beta4 gene, alpha6 gene, or laminin5-gamma2 subunit gene, for example, primer or probe or a primer or probe for any of the sequences or fragments of the sequences, set forth in SEQ ID NOs:1,3,5,7,9,11, and 13 can be less than or equal to about 6, 7, 8, 9, 10, 11, 12 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 125, 150, 175, 200, 225, 250, 275, 300, 325, 350, 375, 400, 425, 450, 475, 500, 550, 600, 650, 700, 750, 800, 850, 900, 950, 1000, 1250, 1500, 1750, 2000, 2250, 2500, 2750, 3000, 3500, or 4000 nucleotides long.

The primers for the beta4 gene, alpha6 gene, or laminin5-gamma2 subunit gene, or any of the sequences or fragments of the sequences, set forth in SEQ ID NOs:1,3,5,7,9,11, and 13, for example, typically will be used to produce an amplified DNA product that contains a desired region. In general, typically the size of the product will be such that the size can be accurately determined to within 3, or 2 or 1 nucleotides.

In certain embodiments this product is at least about 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 125, 150, 175, 200, 225, 250, 275, 300, 325, 350, 375, 400, 425, 450, 475, 500, 550, 600, 650, 700, 750, 800, 850, 900, 950, 1000, 1250, 1500, 1750, 2000, 2250, 2500, 2750, 3000, 3500, or 4000 nucleotides long.

In other embodiments the product is less than or equal to about 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 125, 150, 175, 200, 225, 250, 275, 300, 325, 350, 375, 400, 425, 450, 475, 500, 550, 600, 650, 700, 750, 800, 850,

900, 950, 1000, 1250, 1500, 1750, 2000, 2250, 2500, 2750, 3000, 3500, or 4000 nucleotides long.

d) Sequence similarities

It is understood that as discussed herein the use of the terms homology and identity mean the same thing as similarity. Thus, for example, if the word homology is used between two non-natural sequences it is understood that this is not necessarily indicating an evolutionary relationship between these two sequences, but rather is looking at the similarity or relatedness between their nucleic acid sequences. Many of the methods for determining homology between two evolutionarily related molecules are routinely applied to any two or more nucleic acids or proteins for the purpose of measuring sequence similarity regardless of whether or not they are evolutionarily related.

In general, it is understood that one way to define any known variants and derivatives or those that might arise, of the disclosed genes and proteins herein, is through defining the variants and derivatives in terms of homology to specific known sequences. This identity of particular sequences disclosed herein is also discussed elsewhere herein. In general, variants of genes and proteins herein disclosed typically have at least, about 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, or 99 percent homology to the stated sequence or the native sequence. Those of skill in the art readily understand how to determine the homology of two proteins or nucleic acids, such as genes. For example, the homology can be calculated after aligning the two sequences so that the homology is at its highest level.

Another way of calculating homology can be performed by published algorithms. Optimal alignment of sequences for comparison may be conducted by the local homology algorithm of Smith and Waterman Adv. Appl. Math. 2: 482 (1981), by the homology alignment algorithm of Needleman and Wunsch, J. MoL Biol. 48: 443 (1970), by the search for similarity method of Pearson and Lipman, Proc. Natl. Acad. Sci. U.S.A. 85: 2444 (1988), by computerized implementations of these algorithms (GAP, BESTFIT, FASTA, and TFASTA in the Wisconsin Genetics Software Package, Genetics Computer Group, 575 Science Dr., Madison, WI), or by inspection.

The same types of homology can be obtained for nucleic acids by for example the

30 algorithms disclosed in Zuker, M. Science 244:48-52, 1989, Jaeger et al. Proc. Natl. Acad. Sci.

USA 86:7706-7710, 1989, Jaeger et al. Methods Enzymol. 183:281-306, 1989 which are herein incorporated by reference for at least material related to nucleic acid alignment. It is understood that any of the methods typically can be used and that in certain instances the results of these various methods may differ, but the skilled artisan understands if identity is found with at least one

35 of these methods, the sequences would be said to have the stated identity, and be disclosed herein.

For example, as used herein, a sequence recited as having a particular percent homology to another sequence refers to sequences that have the recited homology as calculated by any one or more of the calculation methods described above. For example, a first sequence has 80 percent homology, as defined herein, to a second sequence if the first sequence is calculated to have 80 5 percent homology to the second sequence using the Zuker calculation method even if the first sequence does not have 80 percent homology to the second sequence as calculated by any of the other calculation methods. As another example, a first sequence has 80 percent homology, as defined herein, to a second sequence if the first sequence is calculated to have 80 percent homology to the second sequence using both the Zuker calculation method and the Pearson and 10 Lipman calculation method even if the first sequence does not have 80 percent homology to the second sequence as calculated by the Smith and Waterman calculation method, the Needleman and Wunsch calculation method, the Jaeger calculation methods, or any of the other calculation methods. As yet another example, a first sequence has 80 percent homology, as defined herein, to a second sequence if the first sequence is calculated to have 80 percent homology to the second 15 sequence using each of calculation methods (although, in practice, the different calculation. methods will often result in different calculated homology percentages).

It is also understood that functional fragments as well as antigenic fragments as well as fragments that can be used in selection protocols of the disclosed compositions are also disclosed. For example, integrins have domains that interact with the other integrins. It may be advantageous in certain embodiments to utilize just the integrin binding domain fragment of, for example, the beta4 integrin, in a selection protocol disclosed herein. By using this domain of the beta4 integrin as the selection target, for example, the selection protocol will be biased for molecules binding this domain of beta4 integrin.

e) Hybridization/selective hybridization

The term hybridization typically means a sequence driven interaction between at least two nucleic acid molecules, such as a primer or a probe and a gene. Sequence driven interaction means an interaction that occurs between two nucleotides or nucleotide analogs or nucleotide derivatives in a nucleotide specific manner. For example, G interacting with C or A interacting with T are sequence driven interactions. Typically sequence driven interactions occur on the Watson-Crick face or Hoogsteen face of the nucleotide. The hybridization of two nucleic acids is affected by a number of conditions and parameters known to those of skill in the art. For example, the salt concentrations, pH, and temperature of the reaction all affect whether two nucleic acid molecules will hybridize.

Parameters for selective hybridization between two nucleic acid molecules are well known to those of skill in the art. For example, in some embodiments selective hybridization conditions

can be defined as stringent hybridization conditions. For example, stringency of hybridization is controlled by both temperature and salt concentration of either or both of the hybridization and washing steps. For example, the conditions of hybridization to achieve selective hybridization may involve hybridization in high ionic strength solution (6X SSC or 6X SSPE) at a temperature 5 that is about 12-25°C below the Tm (the melting temperature at which half of the molecules dissociate from their hybridization partners) followed by washing at a combination of temperature and salt concentration chosen so that the washing temperature is about 5°C to 20°C below the Tm. The temperature and salt conditions are readily determined empirically in preliminary experiments in which samples of reference DNA immobilized on filters are hybridized to a labeled 10 nucleic acid of interest and then washed under conditions of different stringencies. Hybridization temperatures are typically higher for DNA-RNA and RNA-RNA hybridizations. The conditions can be used as described above to achieve stringency, or as is known in the art. (Sambrook et al., Molecular Cloning: A Laboratory Manual, 2nd Ed., Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, 1989; Kunkel et al. Methods Enzymol. 1987:154:367, 1987 which is herein 15 incorporated by reference for material at least related to hybridization of nucleic acids). A preferable stringent hybridization condition for a DNA:DNA hybridization can be at about 68°C (in aqueous solution) in 6X SSC or 6X SSPE followed by washing at 68°C. Stringency of hybridization and washing, if desired, can be reduced accordingly as the degree of complementarity desired is decreased, and further, depending upon the G-C or A-T richness of any 20 area wherein variability is searched for. Likewise, stringency of hybridization and washing, if desired, can be increased accordingly as homology desired is increased, and further, depending upon the G-C or A-T richness of any area wherein high homology is desired, all as known in the art.

Another way to define selective hybridization is by looking at the amount (percentage) of one of the nucleic acids bound to the other nucleic acid. For example, in some embodiments selective hybridization conditions would be when at least about, 60, 65, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100 percent of the limiting nucleic acid is bound to the non-limiting nucleic acid. Typically, the non-limiting primer is in for example, 10 or 100 or 1000 fold excess. This type of assay can be performed at under conditions where both the limiting and non-limiting primer are for example, 10 fold or 100 fold or 1000 fold below their k_d, or where only one of the nucleic acid molecules is 10 fold or 100 fold or 1000 fold or where one or both nucleic acid molecules are above their k_d.

Another way to define selective hybridization is by looking at the percentage of primer that gets enzymatically manipulated under conditions where hybridization is required to promote the desired enzymatic manipulation. For example, in some embodiments selective hybridization conditions would be when at least about, 60, 65, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82,

83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100 percent of the primer is enzymatically manipulated under conditions which promote the enzymatic manipulation, for example if the enzymatic manipulation is DNA extension, then selective hybridization conditions would be when at least about 60, 65, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100 percent of the primer molecules are extended. Preferred conditions also include those suggested by the manufacturer or indicated in the art as being appropriate for the enzyme performing the manipulation.

Just as with homology, it is understood that there are a variety of methods herein disclosed for determining the level of hybridization between two nucleic acid molecules. It is understood that these methods and conditions may provide different percentages of hybridization between two nucleic acid molecules, but unless otherwise indicated meeting the parameters of any of the methods would be sufficient. For example if 80% hybridization was required and as long as hybridization occurs within the required parameters in any one of these methods it is considered disclosed herein.

It is understood that those of skill in the art understand that if a composition or method meets any one of these criteria for determining hybridization either collectively or singly it is a composition or method that is disclosed herein.

f) Delivery of the compositions to cells

The disclosed compositions and methods often entail delivery of the compositions to cells.

20 For example, antisense molecules directed to alpha6 mRNA or gamma2 mRNA can be delivered to cells via any method. A number of exemplary methods are disclosed herein. It is also understood that in certain embodiments, non-nucleic acid molecules will be and can be delivered to cells, for example an antibody to beta4 integrin, alpha6 integrin or, gamma2, or a small molecule, or a peptide. Delivery of these molecules can occur by any means, and exemplary compositions and methods for such delivery are disclosed herein.

There are a number of compositions and methods which can be used to deliver nucleic acids to cells, either in vitro or in vivo. These methods and compositions can largely be broken down into two classes: viral based delivery systems and non-viral based delivery systems. For example, the nucleic acids can be delivered through a number of direct delivery systems such as, electroporation, lipofection, calcium phosphate precipitation, plasmids, viral vectors, viral nucleic acids, phage nucleic acids, phages, cosmids, or via transfer of genetic material in cells or carriers such as cationic liposomes. Appropriate means for transfection, including viral vectors, chemical transfectants, or physico-mechanical methods such as electroporation and direct diffusion of DNA, are described by, for example, Wolff, J. A., et al., Science, 247, 1465-1468, (1990); and Wolff, J.

use with the compositions and methods described herein. In certain cases, the methods will be modified to specifically function with large DNA molecules. Further, these methods can be used to target certain diseases and cell populations by using the targeting characteristics of the carrier.

(1) Nucleic acid based delivery systems

Transfer vectors can be any nucleotide construction used to deliver genes into cells (e.g., a plasmid), or as part of a general strategy to deliver genes, e.g., as part of recombinant retrovirus or adenovirus (Ram et al. Cancer Res. 53:83-88, (1993)).

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As used herein, plasmid or viral vectors are agents that transport the disclosed nucleic acids, such as gamma2 antisense producing molecules into the cell without degradation and 10 include a promoter yielding expression of the gene in the cells into which it is delivered. In some embodiments the delivery vectors are derived from either a virus or a retrovirus. Viral vectors are, for example, Adenovirus, Adeno-associated virus, Herpes virus, Vaccinia virus, Polio virus, AIDS virus, neuronal trophic virus, Sindbis and other RNA viruses, including these viruses with the HIV backbone. Also preferred are any viral families which share the properties of these viruses which 15 make them suitable for use as vectors. Retroviruses include Murine Moloney Leukemia virus, MMLV, and retroviruses that express the desirable properties of MMLV as a vector. Retroviral vectors are able to carry a larger genetic payload, i.e., a transgene or marker gene, than other viral vectors, and for this reason are a commonly used vector. However, other than Lentivirus vectors, they are typically not as useful in non-proliferating cells. Adenovirus vectors are relatively stable 20 and easy to work with, have high titers, and can be delivered in aerosol formulation, and can transfect non-dividing cells. Pox viral vectors are large and have several sites for inserting genes, they are thermostable and can be stored at room temperature. A preferred embodiment is a viral vector which has been engineered so as to suppress the immune response of the host organism, elicited by the viral antigens. Preferred vectors of this type will carry coding regions for 25 Interleukin 8 or 10.

Viral vectors can have higher transaction (ability to introduce genes) abilities than chemical or physical methods to introduce genes into cells. Typically, viral vectors contain, nonstructural early genes, structural late genes, an RNA polymerase III transcript, inverted terminal repeats necessary for replication and encapsidation, and promoters to control the transcription and replication of the viral genome. When engineered as vectors, viruses typically have one or more of the early genes removed and a gene or gene/promoter cassette is inserted into the viral genome in place of the removed viral DNA. Constructs of this type can carry up to about 8 kb of foreign genetic material. The necessary functions of the removed early genes are typically supplied by cell lines which have been engineered to express the gene products of the early genes in trans.

(a) Retroviral Vectors

A retrovirus is an animal virus belonging to the virus family of Retroviridae, including any types, subfamilies, genus, or tropisms. Retroviral vectors, in general, are described by Verma, I.M., Retroviral vectors for gene transfer. In Microbiology-1985, American Society for Microbiology, pp. 229-232, Washington, (1985), which is incorporated by reference herein. Examples of methods for using retroviral vectors for gene therapy are described in U.S. Patent Nos. 4,868,116 and 4,980,286; PCT applications WO 90/02806 and WO 89/07136; and Mulligan, (Science 260:926-932 (1993)); the teachings of which are incorporated herein by reference.

A retrovirus is essentially a package which has packed into it nucleic acid cargo. The 10 nucleic acid cargo carries with it a packaging signal, which ensures that the replicated daughter molecules will be efficiently packaged within the package coat. In addition to the package signal, there are a number of molecules which are needed in cis, for the replication, and packaging of the replicated virus. Typically a retroviral genome, contains the gag, pol, and env genes which are involved in the making of the protein coat. It is the gag, pol, and env genes which are typically 15 replaced by the foreign DNA that it is to be transferred to the target cell. Retrovirus vectors typically contain a packaging signal for incorporation into the package coat, a sequence which signals the start of the gag transcription unit, elements necessary for reverse transcription, including a primer binding site to bind the tRNA primer of reverse transcription, terminal repeat sequences that guide the switch of RNA strands during DNA synthesis, a purine rich sequence 5' to 20 the 3' LTR that serve as the priming site for the synthesis of the second strand of DNA synthesis, and specific sequences near the ends of the LTRs that enable the insertion of the DNA state of the retrovirus to insert into the host genome. The removal of the gag, pol, and env genes allows for about 8 kb of foreign sequence to be inserted into the viral genome; become reverse transcribed, and upon replication be packaged into a new retroviral particle. This amount of nucleic acid is 25 sufficient for the delivery of a one to many genes depending on the size of each transcript. It is preferable to include either positive or negative selectable markers along with other genes in the insert.

Since the replication machinery and packaging proteins in most retroviral vectors have been removed (gag, pol, and env), the vectors are typically generated by placing them into a packaging cell line. A packaging cell line is a cell line which has been transfected or transformed with a retrovirus that contains the replication and packaging machinery, but lacks any packaging signal. When the vector carrying the DNA of choice is transfected into these cell lines, the vector containing the gene of interest is replicated and packaged into new retroviral particles, by the machinery provided in cis by the helper cell. The genomes for the machinery are not packaged because they lack the necessary signals.

(b) Adenoviral Vectors

The construction of replication-defective adenoviruses has been described (Berkner et al., J. Virology 61:1213-1220 (1987); Massie et al., Mol. Cell. Biol. 6:2872-2883 (1986); Haj-Ahmad et al., J. Virology 57:267-274 (1986); Davidson et al., J. Virology 61:1226-1239 (1987); 5 Zhang "Generation and identification of recombinant adenovirus by liposome-mediated transfection and PCR analysis" BioTechniques 15:868-872 (1993)). The benefit of the use of these viruses as vectors is that they are limited in the extent to which they can spread to other cell types, since they can replicate within an initial infected cell, but are unable to form new infectious viral particles. Recombinant adenoviruses have been shown to achieve high efficiency gene 10 transfer after direct, in vivo delivery to airway epithelium, hepatocytes, vascular endothelium, CNS parenchyma and a number of other tissue sites (Morsy, J. Clin. Invest. 92:1580-1586 (1993); Kirshenbaum, J. Clin. Invest. 92:381-387 (1993); Roessler, J. Clin. Invest. 92:1085-1092 (1993); Moullier, Nature Genetics 4:154-159 (1993); La Salle, Science 259:988-990 (1993); Gomez-Foix, J. Biol. Chem. 267:25129-25134 (1992); Rich, Human Gene Therapy 4:461-476 15 (1993); Zabner, Nature Genetics 6:75-83 (1994); Guzman, Circulation Research 73:1201-1207 (1993); Bout, Human Gene Therapy 5:3-10 (1994); Zabner, Cell 75:207-216 (1993); Caillaud, Eur. J. Neuroscience 5:1287-1291 (1993); and Ragot, J. Gen. Virology 74:501-507 (1993)). Recombinant adenoviruses achieve gene transduction by binding to specific cell surface receptors, after which the virus is internalized by receptor-mediated endocytosis, in the same manner as wild 20 type or replication-defective adenovirus (Chardonnet and Dales, Virology 40:462-477 (1970); Brown and Burlingham, J. Virology 12:386-396 (1973); Svensson and Persson, J. Virology 55:442-449 (1985); Seth, et al., J. Virol. 51:650-655 (1984); Seth, et al., Mol. Cell. Biol. 4:1528-1533 (1984); Varga et al., J. Virology 65:6061-6070 (1991); Wickham et al., Cell 73:309-319 (1993)).

A viral vector can be one based on an adenovirus which has had the E1 gene removed and these virons are generated in a cell line such as the human 293 cell line. In another preferred embodiment both the E1 and E3 genes are removed from the adenovirus genome.

(c) Adeno-associated viral vectors

Another type of viral vector is based on an adeno-associated virus (AAV). This defective parvovirus is a preferred vector because it can infect many cell types and is nonpathogenic to humans. AAV type vectors can transport about 4 to 5 kb and wild type AAV is known to stably insert into chromosome 19. Vectors which contain this site specific integration property are preferred. An especially preferred embodiment of this type of vector is the P4.1 C vector produced by Avigen, San Francisco, CA, which can contain the herpes simplex virus thymidine kinase gene, HSV-tk, and/or a marker gene, such as the gene encoding the green fluorescent protein, GFP.

In another type of AAV virus, the AAV contains a pair of inverted terminal repeats (ITRs) which flank at least one cassette containing a promoter which directs cell-specific expression operably linked to a heterologous gene. Heterologous in this context refers to any nucleotide sequence or gene which is not native to the AAV or B19 parvovirus.

Typically the AAV and B19 coding regions have been deleted, resulting in a safe, noncytotoxic vector. The AAV ITRs, or modifications thereof, confer infectivity and site-specific integration, but not cytotoxicity, and the promoter directs cell-specific expression. United states Patent No. 6,261,834 is herein incorporated by reference for material related to the AAV vector.

The vectors of the present invention thus provide DNA molecules which are capable of integration into a mammalian chromosome without substantial toxicity.

The inserted genes in viral and retroviral usually contain promoters, and/or enhancers to help control the expression of the desired gene product. A promoter is generally a sequence or sequences of DNA that function when in a relatively fixed location in regard to the transcription start site. A promoter contains core elements required for basic interaction of RNA polymerase and transcription factors, and may contain upstream elements and response elements.

(d) Large payload viral vectors

Molecular genetic experiments with large human herpesviruses have provided a means whereby large heterologous DNA fragments can be cloned, propagated and established in cells permissive for infection with herpesviruses (Sun et al., Nature genetics 8: 33-41, 1994; Cotter and Robertson, Curr Opin Mol Ther 5: 633-644, 1999). These large DNA viruses (herpes simplex virus (HSV) and Epstein-Barr virus (EBV), have the potential to deliver fragments of human heterologous DNA > 150 kb to specific cells. EBV recombinants can maintain large pieces of DNA in the infected B-cells as episomal DNA. Individual clones carried human genomic inserts up to 330 kb appeared genetically stable The maintenance of these episomes requires a specific EBV nuclear protein, EBNA1, constitutively expressed during infection with EBV. Additionally, these vectors can be used for transfection, where large amounts of protein can be generated transiently in vitro. Herpesvirus amplicon systems are also being used to package pieces of DNA > 220 kb and to infect cells that can stably maintain DNA as episomes.

Other useful systems include, for example, replicating and host-restricted non-replicating 30 vaccinia virus vectors.

(2) Non-nucleic acid based systems

The disclosed compositions can be delivered to the target cells in a variety of ways. For example, the compositions can be delivered through electroporation, or through lipofection, or through calcium phosphate precipitation. The delivery mechanism chosen will depend in part on

the type of cell targeted and whether the delivery is occurring for example in vivo or in vitro.

Thus, the compositions can comprise, in addition to the disclosed nucleic acids and proteins or vectors for example, lipids such as liposomes, such as cationic liposomes (e.g., DOTMA, DOPE, DC-cholesterol) or anionic liposomes. Liposomes can further comprise proteins to facilitate targeting a particular cell, if desired. Administration of a composition comprising a compound and a cationic liposome can be administered to the blood afferent to a target organ or inhaled into the respiratory tract to target cells of the respiratory tract. Regarding liposomes, see, e.g., Brigham et al. Am. J. Resp. Cell. Mol. Biol. 1:95-100 (1989); Felgner et al. Proc. Natl. Acad. Sci USA 84:7413-7417 (1987); U.S. Pat. No.4,897,355. Furthermore, the compound can be administered as a component of a microcapsule that can be targeted to specific cell types, such as macrophages, or where the diffusion of the compound or delivery of the compound from the microcapsule is designed for a specific rate or dosage.

In the methods described above which include the administration and uptake of exogenous DNA into the cells of a subject (i.e., gene transduction or transfection), delivery of the compositions to cells can be via a variety of mechanisms. As one example, delivery can be via a liposome, using commercially available liposome preparations such as LIPOFECTIN, LIPOFECTAMINE (GIBCO-BRL, Inc., Gaithersburg, MD), SUPERFECT (Qiagen, Inc. Hilden, Germany) and TRANSFECTAM (Promega Biotec, Inc., Madison, WI), as well as other liposomes developed according to procedures standard in the art. In addition, the nucleic acid or vector of this invention can be delivered *in vivo* by electroporation, the technology for which is available from Genetronics, Inc. (San Diego, CA) as well as by means of a SONOPORATION machine (ImaRx Pharmaceutical Corp., Tucson, AZ).

The materials may be in solution or suspension (for example, incorporated into microparticles, liposomes, or cells). These may be targeted to a particular cell type via antibodies, receptors, or receptor ligands. The following references are examples of the use of this technology to target specific proteins to tumor tissue (Senter, et al., Bioconjugate Chem., 2:447-451, (1991); Bagshawe, K.D., Br. J. Cancer, 60:275-281, (1989); Bagshawe, et al., Br. J. Cancer, 58:700-703, (1988); Senter, et al., Bioconjugate Chem., 4:3-9, (1993); Battelli, et al., Cancer Immunol. Immunother., 35:421-425, (1992); Pietersz and McKenzie, Immunolog. Reviews, 129:57-80, (1992); and Roffler, et al., Biochem. Pharmacol, 42:2062-2065, (1991)). These techniques can be used for a variety of other specific cell types. Vehicles, among others, include "stealth" and other antibody conjugated liposomes (including lipid mediated drug targeting to colonic carcinoma), receptor mediated targeting of DNA through cell specific ligands, lymphocyte directed tumor targeting, and highly specific therapeutic retroviral targeting of murine glioma cells in vivo. The following references are examples of the use of this technology to target specific proteins to tumor

tissue (Hughes et al., Cancer Research, 49:6214-6220, (1989); and Litzinger and Huang,

Biochimica et Biophysica Acta, 1104:179-187, (1992)). In general, receptors are involved in
pathways of endocytosis, either constitutive or ligand induced. These receptors cluster in clathrincoated pits, enter the cell via clathrin-coated vesicles, pass through an acidified endosome in which
the receptors are sorted, and then either recycle to the cell surface, become stored intracellularly,
or are degraded in lysosomes. The internalization pathways serve a variety of functions, such as
nutrient uptake, removal of activated proteins, clearance of macromolecules, opportunistic entry of
viruses and toxins, dissociation and degradation of ligand, and receptor-level regulation. Many
receptors follow more than one intracellular pathway, depending on the cell type, receptor
concentration, type of ligand, ligand valency, and ligand concentration. Molecular and cellular
mechanisms of receptor-mediated endocytosis has been reviewed (Brown and Greene, DNA and
Cell Biology 10:6, 399-409 (1991)).

Nucleic acids that are delivered to cells which are to be integrated into the host cell genome, typically contain integration sequences. These sequences are often viral related

15 sequences, particularly when viral based systems are used. These viral integration systems can also be incorporated into nucleic acids which are to be delivered using a non-nucleic acid based system of deliver, such as a liposome, so that the nucleic acid contained in the delivery system can be come integrated into the host genome.

Other general techniques for integration into the host genome include, for example,

systems designed to promote homologous recombination with the host genome. These systems typically rely on sequence flanking the nucleic acid to be expressed that has enough homology with a target sequence within the host cell genome that recombination between the vector nucleic acid and the target nucleic acid takes place, causing the delivered nucleic acid to be integrated into the host genome. These systems and the methods necessary to promote homologous

recombination are known to those of skill in the art.

(3) In vivo/ex vivo

As described above, the compositions can be administered in a pharmaceutically acceptable carrier and can be delivered to the subject§s cells *in vivo* and/or *ex vivo* by a variety of mechanisms well known in the art (e.g., uptake of naked DNA, liposome fusion, intramuscular injection of DNA via a gene gun, endocytosis and the like).

If ex vivo methods are employed, cells or tissues can be removed and maintained outside the body according to standard protocols well known in the art. The compositions can be introduced into the cells via any gene transfer mechanism, such as, for example, calcium phosphate mediated gene delivery, electroporation, microinjection or proteoliposomes. The transduced cells can then be infused (e.g., in a pharmaceutically acceptable carrier) or

homotopically transplanted back into the subject per standard methods for the cell or tissue type. Standard methods are known for transplantation or infusion of various cells into a subject.

It is understood that in certain embodiments, constructs which produce an integrin signal transduction inhibitor are driven by inducible promoters, rather than constitutive promoters.

5 inducible systems provide certain advantages, to the expression of the disclosed constructs. Any inducible system can be used. Also disclosed are cells containing the inducible systems, described herein, and in the Examples. These cells, can be used as model systems in a wide variety of assays, as well as in vivo settings.

a) Expression systems

The nucleic acids that are delivered to cells typically contain expression controlling systems. For example, the inserted genes in viral and retroviral systems usually contain promoters, and/or enhancers to help control the expression of the desired gene product. A promoter is generally a sequence or sequences of DNA that function when in a relatively fixed location in regard to the transcription start site. A promoter contains core elements required for basic interaction of RNA polymerase and transcription factors, and may contain upstream elements and response elements.

(1) Viral Promoters and Enhancers

Preferred promoters controlling transcription from vectors in mammalian host cells may be obtained from various sources, for example, the genomes of viruses such as: polyoma, Simian Virus 40 (SV40), adenovirus, retroviruses, hepatitis-B virus and most preferably cytomegalovirus, or from heterologous mammalian promoters, e.g. beta actin promoter. The early and late promoters of the SV40 virus are conveniently obtained as an SV40 restriction fragment which also contains the SV40 viral origin of replication (Fiers et al., Nature, 273: 113 (1978)). The immediate early promoter of the human cytomegalovirus is conveniently obtained as a HindIII E restriction fragment (Greenway, P.J. et al., Gene 18: 355-360 (1982)). Of course, promoters from the host cell or related species also are useful herein.

Enhancer generally refers to a sequence of DNA that functions at no fixed distance from the transcription start site and can be either 5' (Laimins, L. et al., Proc. Natl. Acad. Sci. 78: 993 (1981)) or 3' (Lusky, M.L., et al., Mol. Cell Bio. 3: 1108 (1983)) to the transcription unit.

30 Furthermore, enhancers can be within an intron (Banerji, J.L. et al., Cell 33: 729 (1983)) as well as within the coding sequence itself (Osborne, T.F., et al., Mol. Cell Bio. 4: 1293 (1984)). They are usually between 10 and 300 bp in length, and they function in cis. Enhancers function to increase transcription from nearby promoters. Enhancers also often contain response elements that mediate the regulation of transcription. Promoters can also contain response elements that mediate

35 the regulation of transcription. Enhancers often determine the regulation of expression of a gene.

While many enhancer sequences are now known from mammalian genes (globin, elastase, albumin, α-fetoprotein and insulin), typically one will use an enhancer from a eukaryotic cell virus for general expression. Preferred examples are the SV40 enhancer on the late side of the replication origin (bp 100-270), the cytomegalovirus early promoter enhancer, the polyoma 5 enhancer on the late side of the replication origin, and adenovirus enhancers.

The promoter and/or enhancer may be specifically activated either by light or specific chemical events which trigger their function. Systems can be regulated by reagents such as tetracycline and dexamethasone. There are also ways to enhance viral vector gene expression by exposure to irradiation, such as gamma irradiation, or alkylating chemotherapy drugs.

In certain embodiments the promoter and/or enhancer region can act as a constitutive promoter and/or enhancer to maximize expression of the region of the transcription unit to be transcribed. In certain constructs the promoter and/or enhancer region be active in all eukaryotic cell types, even if it is only expressed in a particular type of cell at a particular time. A preferred promoter of this type is the CMV promoter (650 bases). Other preferred promoters are SV40 promoters, cytomegalovirus (full length promoter), and retroviral vector LTF.

It has been shown that all specific regulatory elements can be cloned and used to construct expression vectors that are selectively expressed in specific cell types such as melanoma cells. The glial fibrillary acetic protein (GFAP) promoter has been used to selectively express genes in cells of glial origin.

Expression vectors used in eukaryotic host cells (yeast, fungi, insect, plant, animal, human or nucleated cells) may also contain sequences necessary for the termination of transcription which may affect mRNA expression. These regions are transcribed as polyadenylated segments in the untranslated portion of the mRNA encoding tissue factor protein. The 3' untranslated regions also include transcription termination sites. It is preferred that the transcription unit also contain a polyadenylation region. One benefit of this region is that it increases the likelihood that the transcribed unit will be processed and transported like mRNA. The identification and use of polyadenylation signals in expression constructs is well established. It is preferred that homologous polyadenylation signals be used in the transgene constructs. In certain transcription units, the polyadenylation region is derived from the SV40 early polyadenylation signal and consists of about 400 bases. It is also preferred that the transcribed units contain other standard sequences alone or in combination with the above sequences improve expression from, or stability of, the construct.

(2) Markers

The viral vectors can include nucleic acid sequence encoding a marker product. This

35 marker product is used to determine if the gene has been delivered to the cell and once delivered is

being expressed. Preferred marker genes are the *E. Coli* lacZ gene, which encodes ß-galactosidase, and green fluorescent protein.

In some embodiments the marker may be a selectable marker. Examples of suitable selectable markers for mammalian cells are dihydrofolate reductase (DHFR), thymidine kinase, neomycin, neomycin analog G418, hydromycin, and puromycin. When such selectable markers are successfully transferred into a mammalian host cell, the transformed mammalian host cell can survive if placed under selective pressure. There are two widely used distinct categories of selective regimes. The first category is based on a cell's metabolism and the use of a mutant cell line which lacks the ability to grow independent of a supplemented media. Two examples are:

10 CHO DHFR- cells and mouse LTK- cells. These cells lack the ability to grow without the addition of such nutrients as thymidine or hypoxanthine. Because these cells lack certain genes necessary for a complete nucleotide synthesis pathway, they cannot survive unless the missing nucleotides are provided in a supplemented media. An alternative to supplementing the media is to introduce an intact DHFR or TK gene into cells lacking the respective genes, thus altering their growth requirements. Individual cells which were not transformed with the DHFR or TK gene will not be capable of survival in non-supplemented media.

The second category is dominant selection which refers to a selection scheme used in any cell type and does not require the use of a mutant cell line. These schemes typically use a drug to arrest growth of a host cell. Those cells which have a novel gene would express a protein conveying drug resistance and would survive the selection. Examples of such dominant selection use the drugs neomycin, (Southern P. and Berg, P., J. Molec. Appl. Genet. 1: 327 (1982)), mycophenolic acid, (Mulligan, R.C. and Berg, P. Science 209: 1422 (1980)) or hygromycin, (Sugden, B. et al., Mol. Cell. Biol. 5: 410-413 (1985)). The three examples employ bacterial genes under eukaryotic control to convey resistance to the appropriate drug G418 or neomycin (geneticin), xgpt (mycophenolic acid) or hygromycin, respectively. Others include the neomycin analog G418 and puromycin.

h) Peptides

(1) Protein variants

As discussed herein there are numerous variants of the beta4 integrin protein, alpha6
integrin protein, and gamma2 laminin5 protein, for example, that are known and herein contemplated. In addition, to the known functional homologue variants there are derivatives of the beta4, alpha6, and gamma2, and other disclosed proteins which also function in the disclosed methods and compositions. Protein variants and derivatives are well understood to those of skill in the art and in can involve amino acid sequence modifications. For example, amino acid sequence modifications typically fall into one or more of three classes: substitutional, insertional or

deletional variants. Insertions include amino and/or carboxyl terminal fusions as well as intrasequence insertions of single or multiple amino acid residues. Insertions ordinarily will be smaller insertions than those of amino or carboxyl terminal fusions, for example, on the order of one to four residues. Immunogenic fusion protein derivatives are made by fusing a polypeptide 5 sufficiently large to confer immunogenicity to the target sequence by cross-linking in vitro or by recombinant cell culture transformed with DNA encoding the fusion. Deletions are characterized by the removal of one or more amino acid residues from the protein sequence. Typically, no more than about from 2 to 6 residues are deleted at any one site within the protein molecule. These variants ordinarily are prepared by site specific mutagenesis of nucleotides in the DNA encoding 10 the protein, thereby producing DNA encoding the variant, and thereafter expressing the DNA in recombinant cell culture. Techniques for making substitution mutations at predetermined sites in DNA having a known sequence are well known, for example M13 primer mutagenesis and PCR mutagenesis. Amino acid substitutions are typically of single residues, but can occur at a number of different locations at once; insertions usually will be on the order of about from 1 to 10 amino 15 acid residues; and deletions will range about from 1 to 30 residues. Deletions or insertions preferably are made in adjacent pairs, i.e. a deletion of 2 residues or insertion of 2 residues. Substitutions, deletions, insertions or any combination thereof may be combined to arrive at a final construct. The mutations must not place the sequence out of reading frame and preferably will not create complementary regions that could produce secondary mRNA structure. Substitutional 20 variants are those in which at least one residue has been removed and a different residue inserted in its place. Such substitutions generally are made in accordance with the following Tables 1 and 2 and are referred to as conservative substitutions.

TABLE 1: Amino Acid Abbreviations

Y

Tyr

tyrosine

	tryptophan	Trp		W
	valine	Val		V
5	TABLE 2: Amino	Acid Sul	bstitutio	ons
,	Original Residue	Evennl	ary Con	nservative Substitutions, others are known in the art.
		Lacinpi	ser	institutive Substitutions, surers are
	Ala			_
	Arg		lys, gln	
	Asn		gln; his	iS .
10	Asp		glu	
	Cys		ser	
	Gln		asn, lys	'S
	Glu		asp	
	Gly		pro	
15	His		asn;gln	n
	Ile		leu; val	
	Leu		ile; val	
	Lys		arg; gli	
	Met		Leu; ile	
20	Phe		met; le	
20	Ser		thr	• •
	Thr		ser	
	Trp		tyr	
	Tyr		trp; ph	ne
25	Val		ile len	11

Substantial changes in function or immunological identity are made by selecting substitutions that are less conservative than those in Table 2, i.e., selecting residues that differ more significantly in their effect on maintaining (a) the structure of the polypeptide backbone in the area of the substitution, for example as a sheet or helical conformation, (b) the charge or hydrophobicity of the molecule at the target site or (c) the bulk of the side chain. The substitutions which in general are expected to produce the greatest changes in the protein properties will be those in which (a) a hydrophilic residue, e.g. seryl or threonyl, is substituted for (or by) a hydrophobic residue, e.g. leucyl, isoleucyl, phenylalanyl, valyl or alanyl; (b) a cysteine or proline is substituted for (or by) any other residue; (c) a residue having an electropositive side chain, e.g., glysyl, arginyl, or histidyl, is substituted for (or by) an electronegative residue, e.g., glutamyl or aspartyl; or (d) a residue having a bulky side chain, e.g., phenylalanine, is substituted for (or by) one not having a side chain, e.g., glycine, in this case, (e) by increasing the number of sites for sulfation and/or glycosylation.

For example, the replacement of one amino acid residue with another that is biologically and/or chemically similar is known to those skilled in the art as a conservative substitution. For example, a conservative substitution would be replacing one hydrophobic residue for another, or one polar residue for another. The substitutions include combinations such as, for example, Gly, Ala; Val, Ile, Leu; Asp, Glu; Asn, Gln; Ser, Thr; Lys, Arg; and Phe, Tyr. Such conservatively substituted variations of each explicitly disclosed sequence are included within the mosaic

polypeptides provided herein.

Substitutional or deletional mutagenesis can be employed to insert sites for N-glycosylation (Asn-X-Thr/Ser) or O-glycosylation (Ser or Thr). Deletions of cysteine or other labile residues also may be desirable. Deletions or substitutions of potential proteolysis sites, e.g. Arg, is accomplished for example by deleting one of the basic residues or substituting one by glutaminyl or histidyl residues.

Certain post-translational derivatizations are the result of the action of recombinant host cells on the expressed polypeptide. Glutaminyl and asparaginyl residues are frequently post-translationally deamidated to the corresponding glutamyl and asparyl residues. Alternatively, these residues are deamidated under mildly acidic conditions. Other post-translational modifications include hydroxylation of proline and lysine, phosphorylation of hydroxyl groups of seryl or threonyl residues, methylation of the o-amino groups of lysine, arginine, and histidine side chains (T.E. Creighton, Proteins: Structure and Molecular Properties, W. H. Freeman & Co., San Francisco pp 79-86 [1983]), acetylation of the N-terminal amine and, in some instances, amidation of the C-terminal carboxyl.

It is understood that one way to define the variants and derivatives of the disclosed proteins herein is through defining the variants and derivatives in terms of homology/identity to specific known sequences. For example, SEQ ID NO:SEQ ID NO:5 sets forth a particular sequence of beta4 integrin cDNA and SEQ ID NO:6 sets forth a particular sequence of a beta4 integrin protein. Specifically disclosed are variants of these and other proteins herein disclosed which have at least, 70% or 75% or 80% or 85% or 90% or 95% homology to the stated sequence. Those of skill in the art readily understand how to determine the homology of two proteins. For example, the homology can be calculated after aligning the two sequences so that the homology is at its highest level.

Another way of calculating homology can be performed by published algorithms. Optimal alignment of sequences for comparison may be conducted by the local homology algorithm of Smith and Waterman Adv. Appl. Math. 2: 482 (1981), by the homology alignment algorithm of Needleman and Wunsch, J. MoL Biol. 48: 443 (1970), by the search for similarity method of Pearson and Lipman, Proc. Natl. Acad. Sci. U.S.A. 85: 2444 (1988), by computerized implementations of these algorithms (GAP, BESTFIT, FASTA, and TFASTA in the Wisconsin Genetics Software Package, Genetics Computer Group, 575 Science Dr., Madison, WI), or by inspection.

The same types of homology can be obtained for nucleic acids by for example the algorithms disclosed in Zuker, M. Science 244:48-52, 1989, Jaeger et al. Proc. Natl. Acad. Sci. 35 USA 86:7706-7710, 1989, Jaeger et al. Methods Enzymol. 183:281-306, 1989 which are herein

incorporated by reference for at least material related to nucleic acid alignment.

It is understood that the description of conservative mutations and homology can be combined together in any combination, such as embodiments that have at least 70% homology to a particular sequence wherein the variants are conservative mutations.

As this specification discusses various proteins and protein sequences it is understood that 5 the nucleic acids that can encode those protein sequences are also disclosed. This would include all degenerate sequences related to a specific protein sequence, i.e. all nucleic acids having a sequence that encodes one particular protein sequence as well as all nucleic acids, including degenerate nucleic acids, encoding the disclosed variants and derivatives of the protein sequences. 10 Thus, while each particular nucleic acid sequence may not be written out herein, it is understood that each and every sequence is in fact disclosed and described herein through the disclosed protein sequence. For example, one of the many nucleic acid sequences that can encode the protein sequence set forth in SEQ ID NO:6 is set forth in SEQ ID NO:5. Another nucleic acid sequence that encodes the same protein sequence set forth in SEQ ID NO:6 is set forth in SEQ ID 15 NO:16. In addition, for example, a disclosed conservative derivative of SEQ ID NO:2 is shown in SEQ ID NO: 17, where the valine (V) at position 34 is changed to a isoleucine (I). It is understood that for this mutation all of the nucleic acid sequences that encode this particular derivative of the beta4 integrin are also disclosed including for example SEQ ID NO:18 and SEQ ID NO:19 which set forth two of the degenerate nucleic acid sequences that encode the particular polypeptide set 20 forth in SEQ ID NO:17. It is also understood that while no amino acid sequence indicates what particular DNA sequence encodes that protein within an organism, where particular variants of a disclosed protein are disclosed herein, the known nucleic acid sequence that encodes that protein in the particular organism from which that protein arises is also known and herein disclosed and described.

i) Pharmaceutical carriers/Delivery of pharmaceutical products

As described above, the compositions, for example, compositions that inhibit alpha6 function, beta4 function, or gamma2 function can also be administered *in vivo* in a pharmaceutically acceptable carrier. By "pharmaceutically acceptable" is meant a material that is not biologically or otherwise undesirable, i.e., the material may be administered to a subject, along with the nucleic acid or vector, without causing any undesirable biological effects or interacting in a deleterious manner with any of the other components of the pharmaceutical composition in which it is contained. The carrier would naturally be selected to minimize any degradation of the active ingredient and to minimize any adverse side effects in the subject, as would be well known to one of skill in the art.

The compositions may be administered orally, parenterally (e.g., intravenously), by

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intramuscular injection, by intraperitoneal injection, transdermally, extracorporeally, topically or the like, although topical intranasal administration or administration by inhalant is typically preferred. As used herein, "topical intranasal administration" means delivery of the compositions into the nose and nasal passages through one or both of the nares and can comprise delivery by a 5 spraying mechanism or droplet mechanism, or through aerosolization of the nucleic acid or vector. The latter may be effective when a large number of animals is to be treated simultaneously. Administration of the compositions by inhalant can be through the nose or mouth via delivery by a spraying or droplet mechanism. Delivery can also be directly to any area of the respiratory system (e.g., lungs) via intubation. The exact amount of the compositions required will vary from subject 10 to subject, depending on the species, age, weight and general condition of the subject, the severity of the allergic disorder being treated, the particular nucleic acid or vector used, its mode of administration and the like. Thus, it is not possible to specify an exact amount for every composition. However, an appropriate amount can be determined by one of ordinary skill in the art using only routine experimentation given the teachings herein.

Parenteral administration of the composition, if used, is generally characterized by injection. Injectables can be prepared in conventional forms, either as liquid solutions or suspensions, solid forms suitable for solution of suspension in liquid prior to injection, or as emulsions. A more recently revised approach for parenteral administration involves use of a slow release or sustained release system such that a constant dosage is maintained. See, e.g., U.S. 20 Patent No. 3,610,795, which is incorporated by reference herein.

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The materials may be in solution, suspension (for example, incorporated into microparticles, liposomes, or cells). These may be targeted to a particular cell type via antibodies, receptors, or receptor ligands. The following references are examples of the use of this technology to target specific proteins to tumor tissue (Senter, et al., Bioconjugate Chem., 2:447-451, (1991); 25 Bagshawe, K.D., Br. J. Cancer, 60:275-281, (1989); Bagshawe, et al., Br. J. Cancer, 58:700-703, (1988); Senter, et al., Bioconjugate Chem., 4:3-9, (1993); Battelli, et al., Cancer Immunol. Immunother., 35:421-425, (1992); Pietersz and McKenzie, Immunolog. Reviews, 129:57-80, (1992); and Roffler, et al., Biochem. Pharmacol, 42:2062-2065, (1991)). Vehicles such as "stealth" and other antibody conjugated liposomes (including lipid mediated drug targeting to 30 colonic carcinoma), receptor mediated targeting of DNA through cell specific ligands, lymphocyte directed tumor targeting, and highly specific therapeutic retroviral targeting of murine glioma cells in vivo. The following references are examples of the use of this technology to target specific proteins to tumor tissue (Hughes et al., Cancer Research, 49:6214-6220, (1989); and Litzinger and Huang, Biochimica et Biophysica Acta, 1104:179-187, (1992)). In general, receptors are involved 35 in pathways of endocytosis, either constitutive or ligand induced. These receptors cluster in clathrin-coated pits, enter the cell via clathrin-coated vesicles, pass through an acidified endosome

in which the receptors are sorted, and then either recycle to the cell surface, become stored intracellularly, or are degraded in lysosomes. The internalization pathways serve a variety of functions, such as nutrient uptake, removal of activated proteins, clearance of macromolecules, opportunistic entry of viruses and toxins, dissociation and degradation of ligand, and receptor-level regulation. Many receptors follow more than one intracellular pathway, depending on the cell type, receptor concentration, type of ligand, ligand valency, and ligand concentration. Molecular and cellular mechanisms of receptor-mediated endocytosis has been reviewed (Brown and Greene, DNA and Cell Biology 10:6, 399-409 (1991)).

(1) Pharmaceutically Acceptable Carriers

The compositions, including antibodies, can be used therapeutically in combination with a pharmaceutically acceptable carrier.

Pharmaceutical carriers are known to those skilled in the art. These most typically would be standard carriers for administration of drugs to humans, including solutions such as sterile water, saline, and buffered solutions at physiological pH. The compositions can be administered intramuscularly or subcutaneously. Other compounds will be administered according to standard procedures used by those skilled in the art.

Pharmaceutical compositions may include carriers, thickeners, diluents, buffers, preservatives, surface active agents and the like in addition to the molecule of choice.

Pharmaceutical compositions may also include one or more active ingredients such as antimicrobial agents, anti-inflammatory agents, anesthetics, and the like.

The pharmaceutical composition may be administered in a number of ways depending on whether local or systemic treatment is desired, and on the area to be treated. Administration may be topically (including ophthalmically, vaginally, rectally, intranasally), orally, by inhalation, or parenterally, for example by intravenous drip, subcutaneous, intraperitoneal or intramuscular injection. The disclosed antibodies can be administered intravenously, intraperitoneally, intramuscularly, subcutaneously, intracavity, or transdermally.

Preparations for parenteral administration include sterile aqueous or non-aqueous solutions, suspensions, and emulsions. Examples of non-aqueous solvents are propylene glycol, polyethylene glycol, vegetable oils such as olive oil, and injectable organic esters such as ethyl oleate. Aqueous carriers include water, alcoholic/aqueous solutions, emulsions or suspensions, including saline and buffered media. Parenteral vehicles include sodium chloride solution, Ringer's dextrose, dextrose and sodium chloride, lactated Ringer's, or fixed oils. Intravenous vehicles include fluid and nutrient replenishers, electrolyte replenishers (such as those based on Ringer's dextrose), and the like. Preservatives and other additives may also be present such as, for

example, antimicrobials, anti-oxidants, chelating agents, and inert gases and the like.

Formulations for topical administration may include ointments, lotions, creams, gels, drops, suppositories, sprays, liquids and powders. Conventional pharmaceutical carriers, aqueous, powder or oily bases, thickeners and the like may be necessary or desirable.

Compositions for oral administration include powders or granules, suspensions or solutions in water or non-aqueous media, capsules, sachets, or tablets. Thickeners, flavorings, diluents, emulsifiers, dispersing aids or binders may be desirable.

Some of the compositions may potentially be administered as a pharmaceutically acceptable acid- or base- addition salt, formed by reaction with inorganic acids such as

10 hydrochloric acid, hydrobromic acid, perchloric acid, nitric acid, thiocyanic acid, sulfuric acid, and phosphoric acid, and organic acids such as formic acid, acetic acid, propionic acid, glycolic acid, lactic acid, pyruvic acid, oxalic acid, malonic acid, succinic acid, maleic acid, and fumaric acid, or by reaction with an inorganic base such as sodium hydroxide, ammonium hydroxide, potassium hydroxide, and organic bases such as mono-, di-, trialkyl and aryl amines and substituted

15 ethanolamines.

(2) Therapeutic Uses

The dosage ranges for the administration of the compositions are those large enough to produce the desired effect in which the symptoms disorder are effected. The dosage should not be so large as to cause adverse side effects, such as unwanted cross-reactions, anaphylactic reactions, and the like. Generally, the dosage will vary with the age, condition, sex and extent of the disease in the patient and can be determined by one of skill in the art. The dosage can be adjusted by the individual physician in the event of any counterindications. Dosage can vary, and can be administered in one or more dose administrations daily, for one or several days.

Other compositions which do not have a specific pharmaceutical function, but which may be used for tracking changes within cellular chromosomes or for the delivery of diagnostic tools for example can be delivered in ways similar to those described for the pharmaceutical products.

j) Chips and micro arrays

Disclosed are chips where at least one address is the sequences or part of the sequences set forth in any of the nucleic acid sequences disclosed herein. Also disclosed are chips where at least one address is the sequences or portion of sequences set forth in any of the peptide sequences disclosed herein.

Also disclosed are chips where at least one address is a variant of the sequences or part of the sequences set forth in any of the nucleic acid sequences disclosed herein. Also disclosed are chips where at least one address is a variant of the sequences or portion of sequences set forth in

any of the peptide sequences disclosed herein.

k) Computer readable mediums

It is understood that the disclosed nucleic acids and proteins can be represented as a sequence consisting of the nucleotides of amino acids. There are a variety of ways to display these sequences, for example the nucleotide guanosine can be represented by G or g. Likewise the amino acid valine can be represented by Val or V. Those of skill in the art understand how to display and express any nucleic acid or protein sequence in any of the variety of ways that exist, each of which is considered herein disclosed. Specifically contemplated herein is the display of these sequences on computer readable mediums, such as, commercially available floppy disks, tapes, chips, hard drives, compact disks, and video disks, or other computer readable mediums. Also disclosed are the binary code representations of the disclosed sequences. Those of skill in the art understand what computer readable mediums. Thus, computer readable mediums on which the nucleic acids or protein sequences are recorded, stored, or saved.

Disclosed are computer readable mediums comprising the sequences and information regarding

15 the sequences set forth herein. Also disclosed are computer readable mediums comprising the

sequences and information regarding the sequences set forth herein wherein the sequences do not

include SEO ID NOs: SEQ ID NOs:1-19.

I) Kits

Disclosed herein are kits that are drawn to reagents that can be used in practicing the methods disclosed herein. The kits can include any reagent or combination of reagent discussed herein or that would be understood to be required or beneficial in the practice of the disclosed methods. For example, the kits could include primers to perform the amplification reactions discussed in certain embodiments of the methods, as well as the buffers and enzymes required to use the primers as intended. For example, disclosed is a kit for assessing a subject's risk for acquiring cancer, such as colon cancer, comprising the primers or probes that hybridize to the sequences set forth in SEQ ID NOs: 1, 3, 5, 7, 9, 11, and 13, for example.

m) Compositions with similar functions

It is understood that the compositions disclosed herein have certain functions, such as inhibiting gamma2 function or binding alpha6 integrin or inhibiting beta4 function. Disclosed herein are certain structural requirements for performing the disclosed functions, and it is understood that there are a variety of structures which can perform the same function which are related to the disclosed structures, and that these structures will ultimately achieve the same result, for example stimulation or inhibition alpha6beta4 signaling or interruption of the alpha6beta4 signaling pathway.

2. Methods of making the compositions

The compositions disclosed herein and the compositions necessary to perform the disclosed methods can be made using any method known to those of skill in the art for that particular reagent or compound unless otherwise specifically noted.

a) Nucleic acid synthesis

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For example, the nucleic acids, such as, the oligonucleotides to be used as primers can be made using standard chemical synthesis methods or can be produced using enzymatic methods or any other known method. Such methods can range from standard enzymatic digestion followed by nucleotide fragment isolation (see for example, Sambrook et al., Molecular Cloning: A Laboratory 10 Manual, 2nd Edition (Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y., 1989)

Chapters 5, 6) to purely synthetic methods, for example, by the cyanoethyl phosphoramidite method using a Milligen or Beckman System 1Plus DNA synthesizer (for example, Model 8700 automated synthesizer of Milligen-Biosearch, Burlington, MA or ABI Model 380B). Synthetic methods useful for making oligonucleotides are also described by Ikuta et al., Ann. Rev. Biochem. 53:323-356 (1984), (phosphotriester and phosphite-triester methods), and Narang et al., Methods Enzymol., 65:610-620 (1980), (phosphotriester method). Protein nucleic acid molecules can be made using known methods such as those described by Nielsen et al., Bioconjug. Chem. 5:3-7 (1994).

b) Peptide synthesis

20 One method of producing the disclosed proteins, or fragments of the disclosed proteins, such as a fragment of SEQ ID NO:6, is to link two or more peptides or polypeptides together by protein chemistry techniques. For example, peptides or polypeptides can be chemically synthesized using currently available laboratory equipment using either Fmoc (9fluorenylmethyloxycarbonyl) or Boc (tert -butyloxycarbonoyl) chemistry. (Applied Biosystems, 25 Inc., Foster City, CA). One skilled in the art can readily appreciate that a peptide or polypeptide corresponding to the disclosed proteins, for example, can be synthesized by standard chemical reactions. For example, a peptide or polypeptide can be synthesized and not cleaved from its synthesis resin whereas the other fragment of a peptide or protein can be synthesized and subsequently cleaved from the resin, thereby exposing a terminal group which is functionally 30 blocked on the other fragment. By peptide condensation reactions, these two fragments can be covalently joined via a peptide bond at their carboxyl and amino termini, respectively, to form an antibody, or fragment thereof. (Grant GA (1992) Synthetic Peptides: A User Guide. W.H. Freeman and Co., N.Y. (1992); Bodansky M and Trost B., Ed. (1993) Principles of Peptide Synthesis. Springer-Verlag Inc., NY (which is herein incorporated by reference at least for 35 material related to peptide synthesis). Alternatively, the peptide or polypeptide is independently synthesized in vivo as described herein. Once isolated, these independent peptides or polypeptides

may be linked to form a peptide or fragment thereof via similar peptide condensation reactions.

For example, enzymatic ligation of cloned or synthetic peptide segments allow relatively short peptide fragments to be joined to produce larger peptide fragments, polypeptides or whole protein domains (Abrahmsen L et al., Biochemistry, 30:4151 (1991)). Alternatively, native chemical ligation of synthetic peptides can be utilized to synthetically construct large peptides or polypeptides from shorter peptide fragments. This method consists of a two step chemical reaction (Dawson et al. Synthesis of Proteins by Native Chemical Ligation. Science, 266:776-779 (1994)). The first step is the chemoselective reaction of an unprotected synthetic peptide--thioester with another unprotected peptide segment containing an amino-terminal Cys residue to give a thioester-linked intermediate as the initial covalent product. Without a change in the reaction conditions, this intermediate undergoes spontaneous, rapid intramolecular reaction to form a native peptide bond at the ligation site (Baggiolini M et al. (1992) FEBS Lett. 307:97-101; Clark-Lewis I et al., J.Biol.Chem., 269:16075 (1994); Clark-Lewis I et al., Biochemistry, 30:3128 (1991); Rajarathnam K et al., Biochemistry 33:6623-30 (1994)).

Alternatively, unprotected peptide segments are chemically linked where the bond formed between the peptide segments as a result of the chemical ligation is an unnatural (non-peptide) bond (Schnolzer, M et al. Science, 256:221 (1992)). This technique has been used to synthesize analogs of protein domains as well as large amounts of relatively pure proteins with full biological activity (deLisle Milton RC et al., Techniques in Protein Chemistry IV. Academic Press, New 20 York, pp. 257-267 (1992)).

The disclosed proteins and polypeptides such as that for SEQ ID NO:6, beta4 integrin, can be made using any traditional recombinant biotechnology method. Examples of such methods can be found in Sambrook et al. which is herein incorporated by reference at least for material related to production of proteins and antibodies.

c) Processes for making the compositions

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Disclosed are processes for making the compositions as well as making the intermediates leading to the compositions. For example, disclosed are nucleic acids in SEQ ID NOs:1, 3, 5, 7, 9, 11, and 13. There are a variety of methods that can be used for making these compositions, such as synthetic chemical methods and standard molecular biology methods. It is understood that the methods of making these and the other disclosed compositions are specifically disclosed.

Disclosed are nucleic acid molecules produced by the process comprising linking in an operative way a nucleic acid molecule comprising the sequence set forth in SEQ ID Nos:1, 3, 5, 7, 9, 11, or 13 or a fragment thereof, and a sequence controlling the expression of the nucleic acid.

Also disclosed are nucleic acid molecules produced by the process comprising linking in

an operative way a nucleic acid molecule comprising a sequence having at least 80% identity to a sequence set forth in SEQ ID Nos:1, 3, 5, 7, 9, 11, or 13 or a fragment thereof, and a sequence controlling the expression of the nucleic acid.

Disclosed are nucleic acid molecules produced by the process comprising linking in an operative way a nucleic acid molecule comprising a sequence that hybridizes under stringent hybridization conditions to a nucleic acid molecule comprising the sequence set forth in SEQ ID Nos:1, 3, 5, 7, 9, 11, or 13 or a fragment thereof, and a sequence controlling the expression of the nucleic acid.

Disclosed are nucleic acid molecules produced by the process comprising linking in an operative way a nucleic acid molecule comprising a sequence encoding a peptide set forth in SEQ ID Nos:2, 4, 6, 8, 10, 12, or 14 or a fragment thereof, and a sequence controlling an expression of the nucleic acid molecule.

Disclosed are nucleic acid molecules produced by the process comprising linking in an operative way a nucleic acid molecule comprising a sequence encoding a peptide having 80% identity to a peptide set forth in SEQ ID Nos:2, 4, 6, 8, 10, 12, or 14 or a fragment thereof and a sequence controlling an expression of the nucleic acid molecule.

Disclosed are nucleic acids produced by the process comprising linking in an operative way a nucleic acid molecule comprising a sequence encoding a peptide having 80% identity to a peptide set forth in SEQ ID Nos:2, 4, 6, 8, 10, 12, or 14 or a fragment thereof, wherein any change from the sequences set forth in SEQ ID Nos:2, 4, 6, 8, 10, 12, or 14 or a fragment thereof are conservative changes and a sequence controlling an expression of the nucleic acid molecule.

Disclosed are cells produced by the process of transforming the cell with any of the disclosed nucleic acids. Disclosed are cells produced by the process of transforming the cell with any of the non-naturally occurring disclosed nucleic acids.

Disclosed are any of the disclosed peptides produced by the process of expressing any of the disclosed nucleic acids. Disclosed are any of the non-naturally occurring disclosed peptides produced by the process of expressing any of the disclosed nucleic acids. Disclosed are any of the disclosed peptides produced by the process of expressing any of the non-naturally disclosed nucleic acids.

Disclosed are animals produced by the process of transfecting a cell within the animal with any of the nucleic acid molecules disclosed herein. Disclosed are animals produced by the process of transfecting a cell within the animal any of the nucleic acid molecules disclosed herein, wherein the animal is a mammal. Also disclosed are animals produced by the process of transfecting a cell within the animal any of the nucleic acid molecules disclosed herein, wherein the mammal is

mouse, rat, rabbit, cow, sheep, pig, or primate.

Also disclose are animals produced by the process of adding to the animal any of the cells disclosed herein.

d) Products produced from selection protocols

Also disclosed are methods of obtaining molecules that act as functional regulators of integrin function, integrin receptor function, and functional regulators of signaling pathways related to integrin receptors, in particular integrin alpha6beta4.

Disclosed are methods for isolating molecules that interact with the proteins set forth in SEQ ID Nos:2, 4, 6, 8, 10, 12, or 14 or a fragment thereof comprising, interacting a library of molecules with the proteins set forth in SEQ ID Nos:2, 4, 6, 8, 10, 12, or 14 or a fragment thereof, removing the unbound molecules, and collecting the molecules that are bound to at least one of the proteins set forth in SEQ ID Nos:2, 4, 6, 8, 10, 12, or 14 or a fragment thereof.

3. Methods of using the compositions

a) Methods of using the compositions as research tools

The compositions can be used for example as targets in combinatorial chemistry protocols or other screening protocols to isolate molecules that possess desired functional properties related to A6B4 and A6B1 signaling pathways.

The disclosed compositions can also be used diagnostic tools related to diseases such as cancer, such as colon cancer.

The disclosed compositions can be used as discussed herein as either reagents in micro arrays or as reagents to probe or analyze existing microarrays. The disclosed compositions can be used in any known method for isolating or identifying single nucleotide polymorphisms. The compositions can also be used in any method for determining allelic analysis of for example, beta4 alleles having varying function, particularly allelic analysis as it relates to beta4 signaling and functions. The compositions can also be used in any known method of screening assays, related to chip/micro arrays. The compositions can also be used in any known way of using the computer readable embodiments of the disclosed compositions, for example, to study relatedness or to perform molecular modeling analysis related to the disclosed compositions.

b) Methods for affecting cancer

The disclosed compositions can be used to affect the growth of cancer cells because as disclosed herein, the disclosed relationships are fundamental to the ability of cancer cells to continue growing. The disclosed compositions, such as antisense constructs that will inhibit the production of either alpha6, beta4, or the gamma2 chain of laminin5 reduce the proliferation of cancer cells. In fact, the compositions cannot only reduce the proliferation of the cancer cells, but

the compositions can kill the cancer cells, as shown herein.

It is understood that cancer is caused by a variety of cellular events, of which certain events related to alpha6 integrin (up regulated), beta4 integrin (upregulated) and gamma2 (upregulated) allow the continued viability of cancer cells, and that interference of these upregulated molecules inhibits the growth and kills the cancer cells. However, there are other known events that can cause non-cancerous cells to become oncogenic. For example, Abl, Ras, EGF receptor, ErB-2, APC, beta-catenin, Arf, Mdm2, p53, Rb, Myc are known to be involved in oncogenesis, and some of these molecules (For example, Ras, APC loss, p53 loss) are directly related to the disclosed target signal transduction pathways mediated by the alpha6beta4 receptor.

Just as the presently disclosed compositions can be used as therapeutics targeting the disclosed relationships, so to there are other targets (For example, Abl, ErB-2) for which pharmaceutical compositions have been developed (For example, Glivec, Herceptin, respectively).

It is understood that the disclosed anti-cancer compositions can be used in combination with other anti-cancer compositions. Great benefits can be obtained from using anti-cancer compositions that target different molecules, in the same signal transduction pathway as well as, or in addition to, targeting molecules in different signal transduction pathways than those disclosed herein. Thus, the disclosed compositions which can affect the growth of cancer cells, indeed kill cancer cells, can be used in conjunction with any other chemotherapy, radiation, or any other anti-cancer therapy.

Disclosed are methods of reducing the proliferation of a cancer cell which comprises inhibiting ligand binding to an integrin receptor on the cancer cell, wherein the integrin receptor comprises an integrin.

Also disclosed are methods of reducing the proliferation of a cancer cell which comprises reducing integrin-integrin interaction, integrin receptor clustering interaction, or integrin-nonintegrin protein interaction.

Also disclosed are methods of selectively reducing the proliferation of cancer cells which comprises reducing integrins from interacting with one another, integrins from clustering, or integrins from interacting with other proteins associated with cancer cells.

Disclosed are methods of reducing the proliferation of a cancer cell which comprises reducing the production of an integrin by the cancer cell.

Also disclosed are methods of reducing the proliferation of a cancer cell which comprises reducing the production of an integrin receptor ligand by the cancer cell.

Disclosed are methods of reducing the proliferation of a cancer cell which comprises

interfering with an integrin signaling pathway.

Also disclosed are methods of selectively killing or reducing the proliferation of cancer cells which comprises inhibiting ligand binding to integrin receptors on the cancer cells, wherein the integrin receptor comprises a B4 integrin.

Disclosed are methods of selectively killing or reducing the proliferation of cancer cells which comprises inhibiting ligand binding to integrin receptors on the cancer cells, wherein the ligand comprises a laminin5.

Also disclosed are methods of selectively killing or reducing the proliferation of cancer cells which comprises inhibiting ligand binding to integrin receptors on the cancer cells, wherein the ligand comprises the gamma2 subunit of the laminin L5.

Disclosed are methods of selectively killing or reducing the proliferation of cancer cells which comprises inhibiting ligand binding to integrin receptors on the cancer cells, wherein the integrin receptor comprises an alpha6 integrin.

Further disclosed are methods of selectively killing or reducing the proliferation of cancer cells which comprises preventing integrin receptor subunits from interacting with one another, preventing integrin clustering, or preventing integrin receptor subunits from interacting with other proteins on cancer cells, wherein the integrin receptor comprises a B4 integrin.

Disclosed are methods of selectively killing or reducing the proliferation of cancer cells which comprises reducing the production of laminin by the cancer cells, wherein the laminin comprises laminin5, and/or any of the subunits of laminin5, such as gamma2.

Also disclosed is a method of selectively killing or reducing the proliferation of cancer cells which comprises interfering with an integrin signaling pathway, wherein the integrin signaling pathway comprises a B4 integrin or an alpha6 integrin or a beta1 integrin or a laminin or a laminin5 or the gamma2 subunit of laminin5.

Also dislossed are methods, wherein the integrin receptor comprises integrin B4 and/or wherein the integrin receptor comprises integrin A6, and/or wherein the ligand that binds to the integrin receptor is laminin5, and/or wherein the integrin receptor is A6B4, and/or wherein the ligand comprises laminin5, and/or wherein the ligand comprises the gamma-2 subunit of laminin5, and/or wherein inhibiting ligand binding to an integrin receptor does not occur by using an antisense molecule to A6, and/or wherein inhibiting ligand binding to an integrin receptor comprises contacting a A6 integrin with a composition that inhibits ligand binding, and/or wherein inhibiting ligand binding to an integrin receptor comprises contacting a B4 integrin with a composition that inhibits ligand binding to an integrin receptor comprises contacting a laminin5 with a composition that inhibits ligand binding, and/or

wherein inhibiting ligand binding to an integrin receptor comprises contacting a gamma-2 subunit with a composition that inhibits ligand binding, and/or wherein reducing integrin-integrin interaction, integrin receptor clustering interaction, or integrin-non-integrin protein interaction does not occur by using a B4-delta-cyt, and/or wherein reducing integrin-integrin interaction, 5 integrin receptor clustering interaction, or integrin-non-integrin protein interaction does not occur by using an antisense molecule to A6, and/or wherein reducing integrin-integrin interaction, integrin receptor clustering interaction, or integrin-non-integrin protein interaction comprises contacting a A6 integrin with a composition that inhibits an interaction between the B4 integrin and another integrin or protein molecule, and/or wherein reducing integrin-integrin interaction, 10 integrin receptor clustering interaction, or integrin-non-integrin protein interaction comprises contacting a B4 integrin with a composition that inhibits the interaction between the B4 integrin and another integrin or protein molecule, and/or wherein reducing integrin-integrin interaction, integrin receptor clustering interaction, or integrin-non-integrin protein interaction comprises contacting a laminin5 with a composition that inhibits ligand binding, and/or wherein reducing 15 integrin-integrin interaction, integrin receptor clustering interaction, or integrin-non-integrin protein interaction comprises contacting a gamma2 subunit with a composition that inhibits ligand binding, and/or wherein the non-integrin protein comprises a growth factor receptor, and/or wherein the non-integrin protein comprises a hemi-desomosome junction, and/or wherein the nonintegrin protein comprises a SH2 domain, and/or wherein the non-integrin protein comprises a Shc 20 protein, and/or wherein the non-integrin protein comprises a IRS-1 protein, and/or wherein the non-integrin protein comprises a IRS-2 protein, and/or wherein reducing the production of an integrin does not occur by using an antisense molecule to A6, and/or wherein the production of an integrin is reduced by inhibiting signaling leading to induction of expression of an integrin, and/or wherein reducing the production of an integrin comprises inhibiting alpha6 production, and/or 25 wherein inhibiting alpha6 production further comprises using antisense molecules to alpha6 mRNA, and/or wherein reducing the production of an integrin comprises inhibiting beta4 production, and/or wherein inhibiting beta4 production further comprises using antisense molecules to beta4 mRNA, and/or wherein reducing the production of an integrin receptor ligand comprises inhibiting gamma2 production, and/or wherein reducing the production of an integrin 30 receptor ligand comprises inhibiting laminin production, and/or wherein reducing the production of an integrin receptor ligand comprises inhibiting laminin5 production, and/or wherein interfering with an integrin signaling pathway does not occur by using a B4-delta-cyt, and/or wherein interfering with an integrin signaling pathway does not occur by using an antisense molecule to A6, and/or wherein interfering with an integrin signaling pathway comprises contacting an A6 35 integrin in the cell with a composition that inhibits ligand binding, and/or wherein interfering with

an integrin signaling pathway comprises contacting a B4 integrin with a composition that inhibits

ligand binding, and/or wherein interfering with an integrin signaling pathway comprises contacting a laminin5 with a composition that inhibits ligand binding, and/or wherein interfering with an integrin signaling pathway comprises contacting a gamma2 subunit with a composition that inhibits ligand binding, and/or wherein interfering with an integrin signaling pathway comprises contacting the cancer cell with a molecule that interferes with at least one of talin, paxillin, vinculin, a CAS family protein, CRX, NCK, FAK, ILK, Src, Fyn, Shc, Grb-2, Guaning nucleotide exchange factors, SOS, DOCK 180, Vav, Syk, P-1-3 kinase, AKT, Bad, Bid, Caspase 9, Cdc42, PAK, Rac, Rho, Rho kinase, Ras, Caveolin, Tetraspan, Receptor-type protein tyrosine phosphatase, SHP-2, Alpha-actinin, Filamin, Cytohesin, Beta3-endonexin, ICAP-1, RACK-1, CIB, actin, receptor tyrosine kinase, IRS-1 or IRS-2, and/or wherein interfering with an integrin signaling pathway comprises contacting the cancer cell with an agent that interferes with post-translational modification of integrins, and/or wherein the post translational modification is glycosylation or phosphorylation.

Also disclosed are methods, wherein the integrin comprises an A6 integrin, and/or wherein 15 the integrin comprises a B4 integrin, and/or further comprising reducing a laminin5-integrin interaction, and/or further comprising reducing a laminin5 gamma2 integrin interaction, and/or wherein the cancer cell comprises normal p53, and/or wherein the proliferation of the cancer cells is not dependent on AKT/PKB, and/or wherein reducing the proliferation of the cancer cells is selective, and/or wherein the cancer cell is not an MDA-MB-435 cell, and/or wherein the cancer 20 cell is not an HMT-3522 cell, and/or wherein the cancer cell is not an RKO colon carcinoma line, and/or wherein the cancer cell does not express exogenous B4 integrin, and/or further comprising contacting the cancer cells with a small molecule, peptide, peptide mimetic, or oligonucleotide or synthetic analog thereof, and/or wherein the cancer cells are contacted with dominant-negative beta 4 integrin, and/or wherein the cancer cell is contacted with an antisense molecule, and/or 25 wherein the antisense molecule is linked to a leader sequence which enables translocation across a cell membrane, and/or wherein the leader sequence binds to a cell surface protein which facilitates internalization, and/or wherein the leader sequence is TAT or antennapedia, or fragment thereof, and/or wherein the antisense molecule is an alpha6 RNA antisense molecule, and/or wherein the small molecule peptide, peptide mimetic, or oligonucleotide or synthetic analog thereof is linked 30 to a carrier, and/or wherein the carrier is at least one of a lipidic carrier, charged carrier, retroviral carrier, TAT or fragment thereof, antennapedia or fragment thereof, or polyethylene glycol, and/or further comprising contacting the cancer cell with another agent which modulates cell signaling, a chemotherapeutic drug, or treated with radiation or angiogenesis inhibitor, and/or wherein reducing the proliferation of cancer cell is in vitro, and/or wherein reducing the proliferation of the 35 cancer cell is in vivo, and/or wherein the cancer cell is selected from the group consisting of melanoma, adenoma, lymphoma, myeloma, carcinoma, plasmocytoma, sarcoma, glioma, thyoma,

leukemia, skin cancer, retinal cancer, breast cancer, prostate cancer, colon cancer, esophageal cancer, stomac cancer, pancreas cancer, brain tumors, lung cancer, ovarian cancer, cervical cancer, hepatic cancer, gastrointestinal cancer, and head and neck cancer cells, and/or wherein the cancer cell is killed, and/or wherein the cancer cell expresses a mutated Ras, and/or wherein the cancer cell expresses a mutated Ras and a mutated p53, and/or wherein the cancer cell expresses a mutated Ras and activates the AKT/PKB protein, and/or wherein the cancer cell expresses a mutated Ras, a mutated p53, and activates the AKT/PKB protein, and/or wherein the cancer cell expresses a mutated Ras and mutated APC, and/or wherein the cancer cell expresses a mutated Ras and mutated APC.

Disclosed are methods of reducing the proliferation of a cancer cell in a patient which comprises administering to the patient a composition which inhibits ligand binding to an integrin receptor on the cancer cell, wherein the integrin receptor comprises an integrin.

Also disclosed are methods of reducing the proliferation of a cancer cell in a patient which comprises administering to the patient a composition which reduces integrin-integrin interaction,

15 integrin receptor clustering interaction, or integrin-non-integrin protein interaction.

Also disclosed are methods of reducing the proliferation of a cancer cell in a patient which comprises administering to the patient a composition which reduces the production of an integrin or laminin by the cancer cell.

Further disclosed are methods of reducing the proliferation of a cancer cell in a patient which comprises administering to the patient a composition which interfers with an integrin signaling pathway.

Also disclsoed are methods, wherein the reduction in cancer cell proliferation is selective, and/or wherein the administering is local or systemic, and/or wherein the patient is additionally administered an agent which modulates cell signaling, a chemotherapeutic drug, or treated with radiation or angiogenesis inhibitor, and/or wherein the additional agent is administered serially or in combination, and/or wherein the local administering is direct application to cancer cells, and/or wherein the direct application to cancer cells is performed during surgery, and/or wherein the direct application to cancer cells is performed topically to cancerous tissue, and/or wherein the systemic administration is by subcutaneous, intraperitoneal, intra-arterial, intravenous, or bolus administration, or by application through a catheter or similar apparatus, and/or wherein the systemic administration comprises a long-term release formulation, and/or wherein the systemic administration is by oral administration, and/or wherein the oral administration comprises administering a pill, capsule, tablet, liquid or suspension.

Disclosed are methods of reducing the proliferation of a cancer cell in a patient which

comprises administering to a patient a vector comprising coding sequence for a protein or peptide which inhibits ligand binding to integrin receptors on cancer cells wherein the coding sequence is under the control of a promoter which functions in mammalian cells.

Also disclosed are methods of reducing the proliferation of a cancer cell in a patient which comprises administering to a patient a vector comprising coding sequence for a protein or peptide which prevents integrin receptor subunits from interacting with one another, prevents integrin receptor clustering interaction, or prevents integrin receptor subunits from interacting with other proteins on cancer cells wherein the coding sequence is under the control of a promoter which functions in mammalian cells.

10 Further disclosed are methods of reducing the proliferation of a cancer cell in a patient which comprises administering to a patient a vector comprising coding sequence for a protein or peptide which prevents integrin receptor subunits from interacting with one another, prevents integrin receptor clustering interaction, or prevents integrin receptor subunits from interacting with other proteins in cancer cells wherein the coding sequence is under the control of a promoter which functions in mammalian cells.

Also disclosed are method of reducing the proliferation of a cancer cell in a patient which comprises administering to a patient a vector comprising coding sequence for a protein or peptide which interferes with integrin subunit or laminin production wherein the coding sequence is under the control of a promoter which functions in mammalian cells.

Further disclosed are methods of reducing the proliferation of a cancer cell in a patient which comprises administering to a patient a vector comprising coding sequence for a protein or peptide which interferes with an integrin signaling pathway of cells wherein the coding sequence is under the control of a promoter which functions in mammalian cells.

Also disclosed are methods, wherein the reduction in cancer cell proliferation is selective,
and/or wherein the vector is administered directly to a cancer cell, and/or wherein the vector is
administered directly to a normal cell, and/or wherein the vector is packaged in a viral vector or
liposome, and/or wherein the vector is a retroviral vector, and/or wherein the vector is
administered systemically, and/or wherein the direct administration is by topical application,
and/or wherein the direct administration is by topical application, and/or wherein the direct
administration is performed during surgery, and/or wherein the direct administration is performed
during surgery, and/or wherein the patient is an animal, such as a mammal, mouse, rabbit, primate,
chimp, ape, goriilla, and human, and/or wherein the cancer cells are selected from the group
consisting of melanoma, adenoma, lymphoma, myeloma, carcinoma, plasmocytoma, sarcoma,
glioma, thyoma, leukemia, skin cancer, retinal cancer, breast cancer, prostate cancer, colon cancer,
esophageal cancer, stomac cancer, pancreas cancer, brain tumors, lung cancer, ovarian cancer,

cervical cancer, hepatic cancer, gastrointestinal cancer, and head and neck cancer cells, and/or wherein the patient is additionally administered at least one of another agent which modulates cell signaling, a chemotherapeutic drug, an angiogenesis inhibitor or treated with radiation, and/or wherein the other agent which modifies cell signaling, chemotherapeutic drug, angiogenesis inhibitor or radiation treatment is administered serially or in combination.

c) Methods of gene modification and gene disruption

The disclosed compositions and methods can be used in targeted gene disruption and modification in any animal that can undergo these events. Gene modification and gene disruption refer to the methods, techniques, and compositions that surround the selective removal or alteration of a gene or stretch of chromosome in an animal, such as a mammal, in a way that propagates the modification through the germ line of the mammal. In general, a cell is transformed with a vector which is designed to homologously recombine with a region of a particular chromosome contained within the cell, as for example, described herein. This homologous recombination event can produce a chromosome which has exogenous DNA introduced, for example in frame, with the surrounding DNA. This type of protocol allows for very specific mutations, such as point mutations, to be introduced into the genome contained within the cell. Methods for performing this type of homologous recombination are disclosed herein.

One of the preferred characteristics of performing homologous recombination in

20 mammalian cells is that the cells should be able to be cultured, because the desired recombination event occur at a low frequency.

Once the cell is produced through the methods described herein, an animal can be produced from this cell through either stem cell technology or cloning technology. For example, if the cell into which the nucleic acid was transfected was a stem cell for the organism, then this cell, after transfection and culturing, can be used to produce an organism which will contain the gene modification or disruption in germ line cells, which can then in turn be used to produce another animal that possesses the gene modification or disruption in all of its cells. In other methods for production of an animal containing the gene modification or disruption in all of its cells, cloning technologies can be used. These technologies generally take the nucleus of the transfected cell and either through fusion or replacement fuse the transfected nucleus with an oocyte which can then be manipulated to produce an animal. The advantage of procedures that use cloning instead of ES technology is that cells other than ES cells can be transfected. For example, a fibroblast cell, which is very easy to culture can be used as the cell which is transfected and has a gene modification or disruption event take place, and then cells derived from this cell can be used to clone a whole animal. Conditional knockouts can also be made which will conditionally delete

expression of the desired molecule, for example, an A6 or integrin or the gamma2 chain of laminin5.

d) Methods of diagnosing cancer

Methods of diagnosing cancer using the disclosed information and the disclosed

5 molecules. In particular disclosed are methods of diagnoses that rely on the combined upregulation of both the ligand and the cogante integrin receptor or cognate integrins. It is understood that all of the methods of diagnosis disclosed herein, can be used with any of the disclosed compositions, but they also can be used in conjunction, where for example, both the ligand and the integrin receptor are monitored and correlated with cancer developing.

Disclosed are methods of assessing a subject's risk of developing cancer comprising determining the amount of A6 present in a target cell obtained from the subject, wherein a determination of increased levels of A6 correlates with an increased risk of cancer.

Disclosed are methods of assessing a subject's risk of acquiring cancer comprising determining the amount of B4 present in a target cell obtained from the subject, wherein a determination of increased levels of B4 correlates with an increased risk of cancer.

Disclosed are methods of assessing a subject's risk of acquiring cancer comprising determining the amount of laminin5 present in a target cell obtained from the subject, wherein a determination of increased levels of laminin5 correlates with an increased risk of cancer.

Disclosed are methods of assessing a subject's risk of acquiring cancer comprising

20 determining the amount of gamma2 subunit present in a target cell obtained from the subject,
wherein a determination of increased levels of gamma2 subunit correlates with an increased risk of
cancer.

Also disclosed are methods, further comprising comparing the amount A6 present to the amount in a control cell, and wherein determining the amount of A6 present in the target cell comprises assaying the amount of A6 mRNA in the cell, and or wherein the assaying the amount of mRNA in the cell comprises hybridizing a A6 probe to a sample of the subject's mRNA, and/or wherein the assaying the amount of mRNA in the cell comprises hybridizing a A6 primer to a sample of the subject's mRNA, and/or wherein the assaying the amount of mRNA further comprises performing an nucleic acid amplification reaction involving the primer, and/or wherein the nucleic acid amplification reaction further comprises performing a polymerase chain reaction on the cDNA, and/or wherein determining the amount of A6 present in the target cell comprises assaying the amount of A6 protein in the cell, and/or further comprising comparing the amount B4 present to the amount in a control cell, and/or further comprising comparing the

amount laminin5 present to the amount in a control cell, and/or further comprising comparing the amount gamma2 subunit present to the amount in a control cell.

C. Examples

Throughout this application, various publications are referenced. The disclosures of these 5 publications in their entireties are hereby incorporated by reference into this application in order to more fully describe the state of the art to which this invention pertains. The references disclosed are also individually and specifically incorporated by reference herein for the material contained in them that is discussed in the sentence in which the reference is relied upon.

It will be apparent to those skilled in the art that various modifications and variations can

be made in the present invention without departing from the scope or spirit of the invention. Other
embodiments of the invention will be apparent to those skilled in the art from consideration of the
specification and practice of the invention disclosed herein. It is intended that the specification
and examples be considered as exemplary only, with a true scope and spirit of the invention being
indicated by the following claims.

The following examples are put forth so as to provide those of ordinary skill in the art with a complete disclosure and description of how the compounds, compositions, articles, devices and/or methods claimed herein are made and evaluated, and are intended to be purely exemplary of the invention and are not intended to limit the scope of what the inventors regard as their invention. Efforts have been made to ensure accuracy with respect to numbers (e.g., amounts, temperature, etc.), but some errors and deviations should be accounted for. Unless indicated otherwise, parts are parts by weight, temperature is in °C or is at ambient temperature, and pressure is at or near atmospheric.

1. Example 1 Colonic epithelial cells dependent on alpha6beta4 receptor signal transduction for growth in the absence of ECM

Integrin alpha6 and integrin beta4 and laminin gamma2 chains are essential for tumor cell survival in vivo. The data disclosed herein indicate that interfering with integrin alpha6-mediated signaling, integrin beta 4signaling, and laminin gamma2 signaling can constitute an effective approach to induce programmed cell death in cancer cells without damaging normal cells.

Ablation of integrin alpha6-dependent signaling, integrin beta4-dependent signaling, and laminin gamma2- dependent signaling in human cancer cell lines supports this.

The murine colonic epithelial transformation system introduced by D'Abaco et al. (1996) was used. In this system, control cells (control), cells containing an activated ras oncogene (Ras), a deletion in the APC gene (APCmin) or both alterations together (Ras+APCmin) can be compared with regard to their proliferation characteristics in tissue culture. The cells were derived from transgenic mice containing a temperature-sensitive allele of SV40 large T under the control of a

gamma interferon-inducible promoter permitting conditional immortalization (Jat et al., 1991).

All experiments were carried out with the cells being kept at non-permissive temperature in the absence of gamma interferon. As shown previously (D'Abaco et al., 1996), only the cells carrying activated Ras and the APCmin mutation were able to form colonies in soft agar in the absence of anchorage to a substratum. All other cell populations did not give rise to colonies under these conditions.

The frequency of tunel-staining cells (Fig.2A) and caspase 3 activity (Fig. 2B) in control, Ras, APCmin and Ras+APCmin colonic epithelial cells (D'Abaco et al., 1996) in suspension was measured. Both assays show a strong suppression of apoptosis in Ras-APCmin cells, indicating that Ras and APCmin co-operate in preventing cell death in the absence of apparent ECM contacts. Activated Ras can protect cells from apoptosis via activation of the serine-threonine kinase AKT (Kauffmann-Zeh et al., 1997; Khwaja et al., 1997). However, both Ras and Ras+APCmin cells show equivalent levels of AKT phosphorylation (not shown), eliminating AKT as the relevant target of Ras+APCmin cooperation.

Although cancer cells have been thought to survive independently of integrin signaling, they frequently express high levels of alpha3 or alpha6 integrin receptors and/or their ligands (Dedhar et al., 1993; Kennel et al., 1989; Koshikawa et al., 1999; Lohi et al., 2000; Nejjari et al., 1999; Van Waes and Carey, 1992). These integrins are suspected to play a role in invasion and metastasis (Mukhopadhyay et al., 1999; Shaw et al., 1997). Ras and APCmin mutations would induce alterations in the mRNA expression patterns of integrin and ECM components. Induction of alpha6 integrin alone, was specific to Ras+APCmin cells (Fig. 3A). In addition, we found elevated expression of the laminin alpha5, gamma2 and beta2 chains induced in Ras and Ras+APCmin cells (Fig. 3B). These chains are components of laminins 5, 10 and 11 (Malinda and Kleinman, 1996), which are ligands of integrin receptors alpha6/beta4 and alpha3/beta1(Kikkawa et al., 2000; Niessen et al., 1994). These results indicate that Ras and APCmin mutations can cooperate to induce autocrine activation of integrin receptors.

Alpha6 integrin can form functional receptors together with either beta1 or beta4 integrins (Clark and Brugge, 1995), leading to signaling through downstream effectors such as focal adhesion kinase (FAK) or the SH2 domain adapter Shc (Giancotti and Ruoslahti, 1999).

30 Conversely, beta4 integrin only binds to alpha6 integrin (Clark and Brugge, 1995). In transformed colonic epithelial cells alpha6 integrin functions in conjunction with integrin beta4 and is engaged by the laminin gamma2 chain to activate Shc (see Fig. 4). Consistent with alpha6 integrin induction, only Ras+APCmin cells can bind a laminin gamma2-specific peptide in an alpha6 and beta4 integrin-specific manner. In contrast, an alpha3/beta1 and alpha6/beta1-specific peptide

35 derived from the laminin gamma1 chain, only binds to control cells (Fig. 4A). In addition, only in

Ras+APCmin cells is She phosphorylated in response to clustering of the integrin alpha6/beta4 receptor by beta4-specific antibodies (Mainiero et al., 1995) and by the laminin gamma2-specific peptide (Fig. 4B). In both cases p52^{Shc} is the major phosphorylated form. Moreover, alpha6 integrin is required for Shc activation, as Shc phosphorylation is inhibited in response to expression of a dominant-negative beta4 integrin mutant. This mutant lacks the cytoplasmic signaling domain but selectively binds to alpha6 integrin to form a ligand binding, yet signaling-defective alpha6/beta4 integrin (Spinardi et al., 1993) (Fig. 4B)

Anti-sense RNAs specific for alpha6 integrin and gamma2 laminin, as well as dominantnegative beta4 integrin, were expressed and results indicated that integrin alpha6, integrin beta4, 10 and laminin gamma2 expression are relevant for the survival of Ras+APCmin cells in Ras+APCmin, Ras, APC and control cells. All three constructs efficiently inhibit the growth of Ras+APCmin cells in soft agar (not shown) and when attached to plastic (Fig. 5A1). In contrast, the proliferation of wt, Ras and APCmin cells is not affected by this treatment (Fig.5A1). Importantly, the colony formation of Ras+APC cells exposed to any of the three inhibitory 15 constructs can be rescued by co-expression of baculovirus p35, a potent inhibitor of caspase activity and of apoptosis (Resnicoff et al., 1998) (Fig. 5A1). Similarly, co-expression of exogenous integrin alpha6 mRNA efficiently rescues Ras-APCmin cell proliferation inhibited by alpha6 anti-sense RNA (Fig. 5A1). This is mirrored by a rescue of alpha6 integrin expression on the cell surface of alpha6 anti-sense RNA expressing cells (Fig. 5B). Moreover, ectopic alpha6 20 integrin expression also rescues apoptosis induced by dominant-negative beta4 integrin (Fig. 5A1). The latter is expressed at constant levels (Fig. 5B). As one would expect, the cells expressing laminin gamma2 antisense RNA could not be rescued by integrin alpha6 overexpression, but showed significant rescue when plated on dishes coated with laminin gamma2 peptide (Fig. 5A1). Similarly, this peptide increases the survival of Ras+APCmin cells in 25 suspension, as indicated by lower caspase 3 activity (Fig. 2C). In summary, the survival of Ras-APCmin cells depends on the expression of alpha6/beta4 integrin and the laminin gamma2 chain. Conversely, the inhibition of these gene activities leads to selective killing of transformed cells.

Data also indicate that cell death due to lack of alpha6/beta4 integrin receptor signaling in Ras-APCmin cells is independent of the death signaling pathway involving the induction of mitochondrial damage and caspase 9 activity. Instead, the data indicate an involvement of the Fas/TNF receptor/death domain protein/caspase8 pathway (Kruidering and Evan, 2000) in the control of tumor cell survival by alpha6 integrin-containing integrin receptors.

Control, APCmin and Ras cells, which lack alpha6/beta4 integrin receptor signaling activity, show high levels of caspase 8 activity when kept in suspension, while at the same time 35 caspase 9 activity cannot be detected (Fig. 2D).

Although Ras-APCmin cells can be rescued from apoptosis by caspase inhibitor baculovirus p35 (Fig. 5 A1, A2 and text above), cell death induced by expression of dominant-negative beta4 integrin cannot be prevented by expression of the survival factor Bcl2 (see Fig. 5 A2). Bcl2 binds to and neutralizes BH3-domain killer proteins that cause mitochondrial damage, cytochrome C release and caspase 9 activation (Luo et al., 1998).

Ras cells which lack alpha6/beta4 integrin receptor signaling activity and cannot survive in the absence of ECM contacts, and Ras+APCmin cells which depend on alpha6/beta4 integrin receptor signaling for survival, show equivalent levels of AKT phosphorylation (data not shown). This indicates that AKT does not serve as the key target for alpha6/beta4 integrin receptor signaling. AKT has been described to promote cell survival via phosphorylation and inactivation of Bad, a BH3-domain killer protein (Datta et al., 1999).

2. Example 2 Other cancer cells dependent on alpha6beta4 receptor signal transduction for growth in the absence of ECM

The survival of highly transformed primary mouse embryo fibroblasts expressing Ras and Myc oncoproteins in conjunction with a homozygous ARF null mutation (Kamijo et al., 1997) depends on expression and function of alpha6 integrin (Fig. 5C), thus demonstrating that this principle also may apply to transformed mesenchymal cells. Moreover, the human colon cancer cell line SW480 carrying multiple oncogenic mutations such as activated Ras (Fujita et al., 1988), amplified c-Myc (Cherif et al., 1988), a mutated APC allele (Munemitsu et al., 1995) and a p53 mutation (Abarzua et al., 1995) is effectively killed by anti-sense integrin alpha6 and laminin gamma2 RNAs as well as dominant-negative beta4 integrin (Fig. 4D). Similarly, integrin alpha6 ablation even leads to apoptosis of Ras/APCmin cells in the presence of active SV40 large T (not shown). Thus the survival of different types of highly transformed cells depends on alpha6 integrin expression, beta4 integrin expression and laminin5 expression irrespective of the status of major tumor supressor genes, such as arf, p53 or rb, and for at least sw480 cells, murine colon epitheilial cells, and the ras/APCmin cells, alpha6, beta4, and laminin5/gamma2 are required. The sensitivity of transformed cells to ablation of integrin signaling, for example, A6B4 integrin receptor signaling, is thus quite remarkable in its apparent generality.

Oncogenic mutations co-operate to engage autocrine integrin signaling. Importantly, this signaling mechanism, involving alpha6 integrin, beta4 integrin and the laminin gamma2 chain, becomes an essential component of the survival mechanism in transformed colonic epithelial cells. In addition, fibroblasts and human colon cancer cells also rely on integrin alpha6 for survival. In contrast, normal and partially transformed cells that express alpha6 integrin at low levels do not require this polypeptide for survival. Thus, integrin signaling inhibition can lead to selective killing of cancer cells.

The laminin-integrin receptor signaling loop is also relevant to the survival of cells transformed by other combinations of oncogenic lesions. Introduction of activated Ras together with dominant-negative p53 (Lloyd et al., 1997) into murine colonic epithelial cells (D'Abaco et al., 1996; see also above) via retroviral infection supports this. Four distinct polyclonal pools of infected cells were derived by drug selection: 1) Control (beo/neo) cells, infected with two retroviruses carrying neomycin (neo) or bleomycin (bleo) drug resistance markers, respectively; 2)Ras cells, infected with a Ras/neo virus and a virus with the bleo marker; 3) DNp53 cells, infected with a DNp53/bleo virus and a virus with the neo marker; and 4) Ras/DNp53 cells, infected with a Ras/neo virus and a DNp53/bleo virus.

As the colon epithelial cells contain temperature-sensitive SV40 large T under control of the gamma interferon promoter, all cell populations were drug-selected at the permissive temperature (33°C) in the presence of gamma interferon. All further experiments were carried out at the non-permissive temperature for SV40 large T in the absence of gamma interferon, as described above for Ras+APCmin cells.

Similar to Ras+APCmin cells, only Ras/DNp53 cells but neither bleo/neo, Ras/bleo or DNp53/neo cells were able to grow in the absence of ECM contact in soft agar (not shown). Moreover, Ras and DNp53 cooperate in the suppression of caspase 3 activity in suspension (Fig. 2B), and in the induction of alpha6 integrin expression (Fig. 4A). In Ras/DNp53 cells passaged through a single round of soft agar growth, alpha6 integrin is expressed at even higher levels, demonstrating a correlation between the ability of the cells to survive in the absence of ECM contact and alpha6 integrin expression. In Ras/DNp53 cells beta4 integrin expression levels are also increased, when compared to controls (Fig. 4A). As expected expression of the laminin gamma2 chain is induced in Ras/bleo and Ras/DNp53 cells (Fig. 4B).

Anti-sense RNAs specific for alpha6 integrin and gamma2 laminin, as well as dominantnegative beta4 integrin were expressed in Ras/DNp53, Ras/bleo, DNp53/neo and neo/bleo cells.
All three constructs efficiently inhibit the growth of Ras/DNp53 cells when attached to plastic. In contrast, the proliferation of neo/bleo, Ras/bleo and DNp53/neo cells is not affected by this treatment (Fig.5A3). Anti-sense alpha6 integrin or dominant-negative beta4 integrin also inhibit the growth of Ras/DNp53 cells in soft agar (Fig. 5A4).

30 Integrin alpha6 expression can also be induced by activated Ras and DNp53 in primary murine colon crypt epithelial cells (not shown). Furthermore introduction of Ras and Myc oncoproteins into our control colon epithelial cells also leads to an induction of alpha6 expression (not shown), suggesting that induction of alpha6 integrin expression may be an integral component of distinct oncogene cooperation paradigms

a) Cell types to be tested

As described herein, there are a variety of cancer and transformed cell types that express alpha6 integrin and laminins at high levels (Dedhar et al., 1993; Koshikawa et al., 1999; Lohi et al., 2000; Van Waes and Carey, 1992). As disclosed herein the following cell lines other than the fibroblasts require alpha6 and beta4 integrin expression, as well as laminin gamma2 chain expression: (1) the human colon carcinoma cell line SW480 (data not shown), (2) Ras+APCmintransformed murine colon epithelial cells (see Fig. 3A) Murine fibroblasts transformed by activated Ras and Myc in conjunction with homozygous Arf null mutation require alpha6 expression for their survival. These cells do not express integrin beta4, but express beta1 integrin, indicating the importance of alpha6/beta1 receptors in the transformed fibroblasts.

b) xxxInducible expression of inhibitors

To interrupt alpha6 integrin and laminin gamma2 chain-dependent signaling in tumors regulatable alpha6 integrin and laminin gamma2 antisense mRNA expression as well as regulatable expression of the beta4 integrin the dominant-negative polypeptide described in

15 Section C were used. Constitutive expression of these inhibitors leads to rapid cell death in tissue culture. In contrast, the establishment of clonal cell lines with inducible expression of anti-sense mRNAs or dominant-negative beta4 integrin minimize such limitations.

The doxycycline (dox)-inducible reverse tetracycline transactivator (rtTA) is frequently used to overexpress transgenes in a temporally regulated fashion in vitro and in vivo (Efrat et al., 1995; Gossen et al., 1995; Ray et al., 1997). These systems are, however, often compromized by the levels of gene expression in the absence of dox administration. The tetracycline controlled transcriptional silencer (tTS), a fusion protein containing the tet repressor and the KRAB-AB domain of the kid-1 transcriptional repressor, is inhibited by doxycycline. As shown in tissue culture (Freundlieb et al., 1999) and in transgenic mice (Zhu et al., 2001), tTS tightens the control of transgene expression in rtTA-based systems, i.e. tTS effectively eliminates leaky baseline expression without altering the inducibility of rtTA-regulated genes. Thus optimal "off/on" regulation of gene expression can be accomplished with the combined use of tTS and rtTA. The complete expression system is commercially available from CLONTECH. This system can be used for the preparation of all inducible cell lines.

Another regulatable system comprising the estrogen-dependent transactivator GalER-VP16 and a promoter under the control of Gal4 DNA binding sites (Braselmann et al., 1993) can be used. (Perez-Roger et al., 1997). This particular experimental set up, was used herein to show that the induction of the beta4 integrin dominant-negative mutant in SW480 colon cancer cells induces apoptosis in vitro, as measured by caspase 3 activation (Fig. 2B). For in vivo use,

35 however, a point mutation has to be introduced into the GalER-VP16 transactivator that eliminates

its sensitivity to estrogen while retaining its response to the anti-estrogen 40H-tamoxifen (Littlewood *et al.*, 1995). 40H-tamoxifen has been demonstrated to reversibly regulate in vivo the activity of another regulatable transactivator, the MycER chimera (Pelengaris *et al.*, 1999).

Typically, inducible cell lines are first stably transfected with the anti-sense or dominantnegative constructs coupled to the regulatable promoters into the four test cell lines. Subsequently, the activator is introduced and the repressor, such as rtTA and tTS (see above) via infection using recombinant retroviruses with different selectable markers

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E. Sequences

1. SEQ ID NO:1 Human alpha6 integrin cds (acc# 4557674)

atggccgccgccggcagctgtgcttgctctacctgtcgggggggctcctgtcccggctcggcgagccttcaacttggac
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 cagcccgaggacaagcggtgttgctcgtgggggccccgcgggagaagcgcttccactgcagagagccaacagaacggga
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 gaaaaaaggcagcatgttaatacgaagcaggaatcccgagacatcttttgggcggtgttatgtcctgagtcagaatctcagg

attgaagacgatatggatgggggagattggagcttttgtgatgggcgattgagaggccatgagaaatttggctcttgccagcaaggtgtagcagctacttttactaaagactttcattacattgtatttggagccccgggtacttataactggaaagggatt gttcgtgtagagcaaaagaataacactttttttgacatgaacatctttgaagatgggccttatgaagttggtggagagact gagcatgatgatgatagtctcgttcctgttcctgctaacagttacttaggtttttctttggactcagggaaaggtattgtttct 5 aaagatgagatcacttttgtatctggtgctcccagagccaatcacagtggagccgtggttttgctgaagagagacatgaag tctgcacatctcctccctgagcacatattcgatggagaaggtctggcctcttcatttggctatgatgtggcggtggtggac ctcaacaaggatgggtggcaagatatagttattggagccccacagtattttgatagagatggagaagttggaggtgcagtg tatgtctacatgaaccagcaaggcagatggaataatgtgaagccaattcgtcttaatggaaccaaagattctatgtttggc attgcagtaaaaaatattggagatattaatcaagatggctacccagatattgcagttggagctccgtatgatgacttggga ggabattcaattgctggaaacatggaccttgatcgaaattcctaccctgatgttgctgttggttccctctcagattcagta actattttcagatcccggcctgtgattaatattcagaaaaccatcacagtaactcctaacagaattgacctccgccagaaa a cag cgtgtgggggcctagtgggatatgcctccaggttaaatcctgttttgaatatactgctaaccccgctggttataat15 aaccaaggttctgagcccaaatatactcaagaactaactctgaagaggcagaaacagaaagtgtgcatggaggaaaccctg tggctacaggataatatcagagataaactgcgtcccattcccataactgcctcagtggagatccaagagccaagctctcgt aggegagtgaattcacttccagaagttcttccaattctgaattcagatgaacccaagacagctcatattgatgttcacttc ttaaaagagggatgtgggagacgacaatgtatgtaacagcaaccttaaactagaatataaattttgcacccgagaaggaaat caagacaaattttcttatttaccaattcaaaaaggtgtaccagaactagttctaaaagatcagaaggatattgctttagaa 20 alaacagtgacaaacagcccttccaacccaaggaatcccacaaaagatggcgatgacgcccatgaggctaaactgattgca acgtttccagacactttaacctattctgcatatagagaactgagggctttccctgagaaacagttgagttgtgttgccaac cagaatggctcgcaagctgactgtgagctcggaaatccttttaaaagaaattcaaatgtcactttttatttggttttaagt acaactgaagtcacctttgacaccccatatctggatattaatctgaagttagaaacaacaagcaatcaagataatttggct ccaattacagctaaagcaaaagtggttattgaactgcttttatcggtctcgggagttgctaaaccttcccaggtgtatttt 25 ggaggtacagttgttggcgagcaagctatgaaatctgaagatgaagtgggaagtttaatagagtatgaattcagggtaata aacttaggtaaacctcttacaaacctcggcacagcaaccttgaacattcagtggccaaaagaaattagcaatgggaaatgg ttgctttatttggtgaaagtagaatccaaaggattggaaaaggtaacttgtgagccacaaaaggagataaactccctgaac gctgaaagaaaataccagactcttaactgtagcgtgaacgtgaactgtgtgaacatcagatgcccgctgcgggggctggac 30 agcaaggcgtctcttattttgcgctcgaggttatggaacagcacatttctagaggaatattccaaactgaactacttggac atteteatgegageetteattgatgtgactgetgetgeegaaatateaggetgeeaaatgeaggeaeteaggttegagtg actgtgtttccctcaaagactgtagctcagtattcgggagtaccttggtggatcatcctagtggctattctcgctgggatc tatcacaaggetgagatccatgetcagecatetgataaagagaggettaettetgatgeatag

2. SEQ ID NO:2 Human integrin alpha6 protein sequence (acc# NP 000201):

1 maaagglcll ylsagllsrl gaafnldtre dnvirkygdp gslfgfslam hwqlqpedkr 61 lllvgaprge alplgranrt gglyscdita rgpctriefd ndadptsesk edgwmgvtvq 121 sqgpggkvvt cahryekrqh vntkqesrdi fgrcyvlsqn lrieddmdgg dwsfcdgrlr 40 181 ghekfgscqq gvaatftkdf hyivfgapgt ynwkgivrve qknntffdmn ifedgpyevg 241 getehdesly pypansylgf sldsgkgivs kdeitfysga pranhsgavy llkrdmksah 301 llpehifdge glassfgydv avvdlnkdgw qdivigapqy fdrdgevgga vyvymnqqgr 361 wnnvkpirln gtkdsmfgia vknigdingd gypdiavgap yddlgkvfiy hgsangintk 421 ptqvlkgisp yfgysiagnm dldrnsypdv avgslsdsvt ifrsrpvini gktitvtpnr 45 481 idlrqktacg apsgiclqvk scfeytanpa gynpsisivg tleaekerrk sglssrvqfr 541 nggsepkytg eltlkrakak vcmeetlwla dnirdklrpi pitasveige pssrrrvnsl 601 pevlpilnsd epktahidvh flkegcgddn vcnsnlkley kfctregnqd kfsylpiqkg 661 vpelvlkdqk dialeitvtn spsnprnptk dgddaheakl iatfpdtlty sayrelrafp 721 ekqlscvang ngsqadcelg npfkrnsnvt fylvlsttev tfdtpyldin lklettsnqd 50 781 nlapitakak vvielllsvs gvakpsqvyf ggtvvgeqam ksedevgsli eyefrvinlg 841 kpltnlgtat lniqwpkeis ngkwllylvk veskglekvt cepqkeinsl nlteshnsrk 901 kreitekqid dnrkfslfae rkyqtlncsv nvncvnircp lrgldskasl ilrsrlwnst 961 fleeyskiny idilmrafid vtaaaeniri pnagtqvrvt vfpsktvaqy sgvpwwiilv 1021 ailagilmla llvfilwkcg ffkrnkkdhy datyhkaeih aqpsdkerlt sda

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3. SEQ ID NO:3 Mus musculus alpha6 integrin cds (acc# 7110658):

atggccgtcgcgggccagttgtgcctgctctacctgtccgcggggcttctagcccggctgggtacagccttcaacctggac
acccgcgaggacaaccgtgatccggaaatcgggggatcccgggagcctcttcggcttctcgccatgcactggcagttg
cagccggaggacaacggctgttgcttgtgggggacctcgggaagcactccggcagaagcactccggcagagggcgaacagagag
ggcctgtacagctgtgacatcacctcccgaggaccttgtacacggattgaatttgataatgacgctgatcctatgtcagaa
agcaaggaagaccagtggatgggagtcactgtccagagccaaggtccagggggcaaaagtggtgacatcgatat
gagaaacggcagcacgtcaacacgaagcagagtcgcgggatatctttgaagatgttatgtcctgagacatccagaa
attgaagatgatatggacggagagctggagttctgcgatggccggttgagaggccatgaaaagttggccctagaaacgcagagcagagcacgtcatttcattacattgttttttggaagccccagggacttacaaccggaagggact
gtccgtgtagaacaaaagaataacactttttttgacatgaacatctttgaagatggccctatgaagttggtggagagaca
gaccatgaaagtctcgtgcccgttcctgctaacagttacctaggcttttcgctggagactcaggaaggcatggagagaca
65 gatcatgatgaaagtctcgtgcccgttcctgctaacagttacctaggcttttcgctggacccaggaaagggaagggaagggatatgttct

aaagatgacatcactbttgtgtctggtgctccaagagccaatcacagtggggctgtagtgttgctaaaaagagacatgaag tccgcacatctgctccctgagtatatattttgacggagaaggcctggcttcctcgtttggctatgatgtggcagtggtggac ctcaatgcagatggtggcaagacatcgttatcggagctccacagtattttgatagggatggtgaagtcgggggtgcagtt tacgtctacattaaccagcaaggcaaatggagtaatgtgaagccgattcgtctaaatggggaccaaagactcgatgtttgga 5 atctctgtgaaaaatataggtgatattaaccaagatggctatccagatattgctgttggagctccctatgatgatctgggg aaggtttttatctatcatggatccccgactggcataattaccaagccaacacggttctcgaggggacatcgccttacttc ggctattcaatcgctgggaatatggacctggatcggaattcctaccccgaccttgctgtgggctccctctcagactcggtc actattttcagatcccggccagtgattaacattctaaaaaccatcacagtgactcctaacagaattgacctccgccagaag tccatgtgtggctcacctagcgggatatgcctcaaggttaaagcctgttttgaatatactgcgaaaccttccggttataac tggctgcaggagaacatcagagacaagctgcgtcccatccccatcacggcttctgtggagatccaggagcccacgtctcgc ttaaaggaaggatgtggagacgacaatgtctgtaacagcaaccttaagctagagtataaatttggtacccgagaaggaaat 15 caagacaaattotottacottocaattoaaaaaggcatoocagaattagtootaaaagatoagaaagatatagototggaa ataacggtgaccaacagcccttcggatccaaggaatccccggaaagatggcgacgatgcccatgaagccaaactcatcgcc acglttccayacactctgacatattccgcttacagagaactgagggctttccctgagaagcagctgagctgtgtggccaac cagaatggctcccaaqccqactqtqaqctcqqaaatcctttcaagagaaattccagtgttactttctatctgattttaagt 20 ccaattacagcgaaggcaaaagtggttattgaattgettttateceteteeggagtegetaageettegeaggtgtatttt ggaggtacagttgttggtgagcaagctatgaaatctgaagatgaagtaggaagtttaatagagtatgaatttagggtgatt aacttaggcaagcctcttaaaaacctcggcacagcaaccttgaatatacagtggcccaaggagattagcaatggcaaatgg 25 cetgaaagaaaataccagaetetcaactgcagegtcaacgtcaggtgtgtgtgaacatcaggtgcccactgcgagggetggac acgaaggcctctctcgttctgtgttccaggttgtggaacagcacatttctagaggaatattccaaactgaactacttggac attetegtgagggettecatagatgteacegetgetgeteagaatateaageteceteaegegggeacteaggttegagtg acqqtqtttccctcaaaqactqtaqctcaqtattcaqqaqtaqcttggtqgatcatcctcctqqctgttcttgccgggatt 30 tatcacaaggctgagatccatactcagccgtctgataaagagaggcttacttccgatgcatag

4. SEQ ID NO:4 Mus musculus alpha6 integrin protein sequence (acc# NP 032423)

1 mavagqlcll ylsagllarl gtafnldtre dnvirksgdp gslfgfslam hwqlqpedkr 61 lllvgaprae alplqranrt gglyscdits rgpctriefd ndadpmsesk edgwmgvtvq 35 121 sqgpggkvvt cahryekrqh vntkqesrdi fgrcyvlsqn lrieddmdgg dwsfcdgrlr 181 ghekfgscqq gvaatftkdf hyivfgapgt ynwkgivrve qknntffdmn ifedgpyevg 241 getdhdeslv pvpansylgf sldsgkgivs kdditfvsga pranhsgavv llkrdmksah 301 llpeyifdge glassfgydv avvdlnadgw qdivigapqy fdrdgevgga vyvyingggk 361 wsnvkpirln gtkdsmfgis vknigdingd gypdiavgap yddlgkvfiy hgsptgiitk 40 421 ptqvlegtsp yfgysiagnm dldrnsypdl avgslsdsvt ifrsrpvini lktitvtpnr 481 idlrqksmcg spsgiclkvk acfeytakps gymppisilg ileaekerrk sglssrvqfr 541 nqgsepkytq eltlnrqkqr acmeetlwlq enirdklrpi pitasveiqe ptsrrrvnsl 601 pevlpilnsn eaktvatovh flkegogddn vonsnlkley kfatregnad kfsylpiaks 661 ipelvlkdak dialeitvtn spsdprnprk daddaheakl iatfattly sayrelrafp 45 721 ekqlscvang ngsqadcelg npfkrnssvt fylilsttev tfdttdldin lklettsngd 781 klapitakak vviellisls gvakpsqvyf ggtvvgeqam ksedevgsli eyefrvinig 841 kplknlgtat lniqwpkeis ngkwllylmk veskglegiv cephneinyl klkeshnsrk 901 krelpekqid darkfalfpe rkyqtlncsv nvrcvnircp lrgldtkasl vlcsrlwnst 961 fleeysklny ldilvrasid vtaaaqnikl phagtqvrvt vfpsktvaqy sgvawwiill 50 1021 avlagilmla llvfllwkcg ffkrnkkdhy datyhkaeih tqpsdkerlt sda

5. SEQ ID NO:5 Human integrin beta4 subunit cds (acc# 6453379):

ggccttcggacagaggtcacctccaagatgttccagaagacgaggactgggtcctttcacatccggcgggggaagtgggt atataccaggtgcagctgcgggcccttgagcacgtggatgggacgcacgtgtgccagctgccggaggaccagaagggcaac atccatctgaaaccttccttctccgacggcctcaagatggacgcgggcatcatctgtgatgtgtgcacctgcgagctgcaa cgtggggagtgccagtgcgggcactgtgtgctacggcgaaggccgctacgagggtcagttctgcgagtatgacaacttc acaggcccaagctgtgactgtcccctcagcaatgccacctgcatcgacagcaatgggggcatctgtaatggacgtggccac tgtgagtgtggccgctgccactgccaccagcagtcgctctacacggacaccatctgcgagatcaactactcggcgtccacc 10 cgggcctctgcgaggacctacgctcctgcgtgcagtgccaggcgtggggcaccggcgagaagaaggggcgcacgtgtgagg aatgcaacttcaaggtcaagatggtggacgagcttaagagaggcgaggaggtggtggtggcgctgctccttccgggacgagg atgacgactgcacctacagctacaccatggaaggtgacggcgccctgggcccaacagcactgtcctggtgcacaagaaga agggactgccctccgggctccttctggtggctcatccccctgctcctcctcctcctgctgctcctggccctgctactgctg ctatgctggaagtactgtgcctgctgcaaggcctgcctggcacttctcccgtgctgcaaccgaggtcacatggtgggcttt 15 aaggaagaccactacatgctgcgggagaacctgatggcctctgaccacttggacacgcccatgctgcgcagcgggaacctc aagggccgtgacgtggtccgctggaaggtcaccaacaacatgcagcggcctggctttgccactcatgccgccagcatcaac cccacagagetggtgccctacgggctgtccttgcgcctggcccgcctttgcaccgagaacctgctgaagcctgacactcgg gagtgcgcccagctgcgccaggaggtggaggagaacctgaacgaggtctacaggcagatctccggtgtacacaagctccag cagaccaagttccggcagcagcccaatgccgggaaaaagcaagaccacaccattgtggacacagtgctgatggcgccccgc 20 tcggccaagccggccctgctgaagcttacagagaagcaggtggaacagagggccttccacgacctcaaggtggcccccggc tactacaccctcactgcagaccaggacgcccggggcatggtggagttccaggagggcgtggagctggtggacgtacgggtg ccctctttatccggcctgaggatgacgacgagaagcagctgctggtggaggccatcgacgtgcccgcaggcactgccacc ctcggccgccgcctggtaaacatcaccatcatcaaggagcaagccagagacgtggtgtcctttgagcagcctgagttctcg gtcagccgcggggaccaggtggcccgcatccctgtcatccggcgtgtcctggacggcgggaagtcccaggtctcctaccgc 25 acacaggatggcaccgcgcagggcaaccgggactacatccccgtggagggtgagctgctgtttccagcctggggaggcctgg aaagagetgeaggtgaageteetggagetgeaagaagttgaeteecteetgeggggeegeeaggteegeegttteeaegte cagctcagcaaccctaagtttggggccacctgggccagcccactccaccaccatcatcatcagggacccagatgaactg gaccggagettcacgagtcagatgttgtcatcacagccacccctcacggcgacctgggcgccccgcagaaccccaatget aaggccgctgggtccaggaagatccatttcaactggctgcccccttctggcaagccaatggggtacagggtaaagtactgg 30 attcagggtgactccgaatccgaagcccacctgctcgacagcaaggtgccctcagtggagctcaccaacctgtacccgtat tgcgactatgagatgaaggtgtgcgcctacggggctcagggcgagggaccctacagctccctggtgtcctgccgcacccac caggaagtgcccagcgagccagggcgtctggccttcaatgtcgtctcctccacggtgacccagctgagctgggctgagccg gctgagaccaacggtgagatcacagcctacgaggtctgctatggcctggtcaacgatgacaaccgacctattggggcccatg aagaaagtgctggttgacaaccctaagaaccggatgctgcttattgagaaccttcgggagtcccagccctaccgctacacg 35 gtgaaggcgcgccaccgggggcctgagcgggaggccatcatcaacctggccacccagcccaagaggcccatg tccatccccatcatccctgacatccctatcgtggacgcccagagcggggaggactacgacagcttccttatgtacagcgat gacgttctacgctctccatcgggcagccagaggcccagcgtctccgatgacactggctgcggctggaagttcgagcccctg ctggggaggagctggacctgcggcgcgtcacgtggcggctgccccgggagctcatcccgcgcctgtcggccagcagcggg 40 gcgacacccgggccccccggagagcacctggtgaatggccggatggactttgccttcccgggcagcaccaactccctgcac aggatgaccacgaccagtgctgctgctatggcacccacctgagcccacacgtgccccaccgcgtgctaagcacatcctcc accetcacacgggactacaactcactgacccgctcagaacactcacactcgaccacactgccgagggactactccaccetc acctccgtctcccaccggcctccctcccatctgggaacacgggaggagcaggcttccgctgtcctgggccctggggtcc45 ccctcctggggcccagactctcgcctgactgctggtgtgcccgacacgcccaccegcctggtgttctctgccctggggccc acatctctcagagtgagctggcaggagccgcggtgcgagcggctgctgcagggctacagtgtggagtaccagctgctgaac ggcggtgagctgcatcggctcaacatccccaaccctgcccagacctcggtggtggtggaagacctcctgccaaccactcc tacgtgttccgcgtgcgggcccagagccaggaaggctggggccgagagcgtgagggtgtcatcaccattgaatcccaggtg cacccgcagagcccactgtgtcccctgccaggctccgccttcactttgagcactcccagtgccccaggcccgctggtgttc 50 actgccctgagcccagactcgctgcagctgagctgggagcggccaccggaggcccaatggggatatcgtcggctacctggtg ccgggcctcagcgagaacgtgccctacaagttcaaggtgcaggccaggaccactgagggcttcgggccagagcgcgagggc atcatcaccatagagtcccaggatggaggtcccttcccgcagctgggcagccgtgccgggctcttccagcacccgctgcaa agcgagtacagcagcatctccaccaccaccagcgccaccgagcccttcctagtgggtccgaccctgggggcccagcac 55 ctggaggcaggctccctcacccggcatgtgacccaggagtttgtgagccggacactgaccaccagcggaacccttagc acccacatggaccaacagttcttccaaacttga

6. SEQ ID NO:6 Human beta4 integrin protein sequence (acc# CAB61345):

1 magprpspwa rlllaalisv slsgtlnrck kapvksctec vrvdkdcayc tdemfrdrrc
61 ntgaellaag cqresivvme ssfqiteetq idttlrrsqm spqglrvrlr pgeerhfele
60 121 vfeplespvd lyilmdfsns msddldnlkk mgqnlarvls qltsdytigf gkfvdkvsvp
181 qtdmrpeklk epwpnsdppf sfknvislte dvdefrnklq gerisgnlda peggfdailq
241 tavctrdigw rpdsthllvf stesafhyea dganvlagim srnderchld ttgtytqyrt
301 qdypsvptlv rllakhniip ifavtnysys yyeklhtyfp vsslgvlqed ssnivellee
361 afnrirsnld iraldsprgl rtevtskmfq ktrtgsfhir rgevgiyqvq lralehvdgt
65 421 hvcqlpedqk gnihlkpsfs dglkmdagii cdvctcelqk evrsarcsfn gdfvcgqcvc
481 segwsgqtcn cstgslsdiq pclregedkp csgrgecqcg hcvcygegry eggfceydnf

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541 qcprtsgflc ndrgrcsmgq cvcepgwtgp scdcplsnat cidsnggicn grghcecgrc
   601 hchqqslytd ticeinysas trasartyap acsarrgapa rrrgarvrna tsrsrwwtsl
   661 rearrwwcaa psgtrmttap tatpwkvtap lgptalswct rrrdcppgsf wwlip11111
   721 lpllallli cwkycacka clallpccnr qhmvqfkedh ymlrenimas dhldtpmlrs
 5 781 gnlkgrdvvr wkvtnnmqrp gfathaasin ptelvpygls lrlarlcten llkpdtreca
   841 qlrqeveenl nevyrqisgv hklqqtkfrq qpnagkkqdh tivdtvlmap rsakpallkl
   901 tekqveqraf hdlkvapgyy tltadqdarg mvefqegvel vdvrvplfir pedddekqll
   961 veaidvpagt atlgrrlvni tiikegardv vsfegpefsv srgdqvarip virrvldggk
1021 sqvsyrtqdg taggnrdyip vegellfqpg eawkelqvkl lelgevdsll rgrqvrrfhv
10 1081 qlsnpkfgah lgqphsttii irdpdeldrs ftsqmlssqp pphgdlgapq npnakaagsr
   1141 kihfnwlpps gkpmgyrvky wiggdsesea hlldskvpsv eltnlypycd yemkvcayga
   1201 qgegpysslv scrthqevps epgrlafnvv sstvtqlswa epaetngeit ayevcyglvn
   1261 ddnrpigpmk kvlvdnpknr mllienlres qpyrytvkar ngagwgpere aiinlatqpk
   1321 rpmsipiipd ipivdaqsge dydsflmysd dvlrspsgsq rpsvsddtgc gwkfepllge
15 1381 eldlrrvtwr lppeliprls assgrssdae aphgppddgg aggkggslpr satpgppgeh
   1441 lvngrmdfaf pgstnslhrm tttsaaaygt hlsphvphrv lstsstltrd ynsltrsehs
   1501 hsttlprdys tltsvsshgl ppiwehgrsr lplswalgsr sraqmkgfpp srgprdsiil
   1561 agrpaapswg pdsrltagvp dtptrlvfsa lgptslrvsw qeprcerplq gysveyqlln
   1621 ggelhrlnip npaqtsvvve dllpnhsyvf rvraqsqegw greregviti esqvhpqspl
20 1681 cplpgsaftl stpsapgplv ftalspdslq lswerprrpn gdivgylvtc emaggggpat
   1741 afrydgdspe srltypglse nypykfkyga rttegfgper egiitiesqd ggpfpqlgsr
   1801 aglfqhplqs eyssisttht satepflvgp tlgaqhleag gsltrhvtqe fvsrtlttsg
   1861 tlsthmdqqf fqt
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25

7. SEQ ID NO:7 Mouse integrin beta4 subunit mRNA (not only cds) (acc#L04678):

ggaccgtcgaggcagcgggactgacccagctgggctcactgtattaagaagcggacccgggacccggagcgcccggggacc 30 tgctgctggcacgaatgctgagtgccagcctccctggagacctggccaaccgctgcaagaaggctcaggtgaagagctgta ccgagtgcatccgggtggacaagagctgtgcctactgcacagacgagctgttcaaggagagggctgcaacacccaggegg acgttctggctgcaggctgcaggggagagagcatcctggtcatggagagcatccttgaaatcacagagaacacccagateg tcaccagcctgcaccgcagccaggtatctccccaaggcctgcaagtccggctgcggcggggtgaggagcgcacgtttgtgt tccaggtctttgagcccctggagagccccgtggatctgtatatcctcatggacttctccaactccatgtctgacgatctgg 35 acaaceteaageagatggggcagaacetggccaagateetgcgccageteaceagegaetacaceattggatttggaaagt ttgtggacaaagtcagcgtcccacagacagacatgaggcccgagaaactgaaggagccctggcccaacagtgatcccccgt tctccttcaaqaacqttatcaqcttaacqqaqaatqtqqaaqaattctqqaacaaactgcaaggagaacgcatctcaggca acctggacgctcctgaagggggctttgatgccatcctgcagacagctgtgtgcacaagggacattggctggagggctgaca gcacccacctgctggtgttctccaccgagtctgccttccactacgaggctgatggtgccaacgttctggccggcatcatga 40 accgcaatgatgagaaatgccacctggacgcctcgggcgcctacacccaatacaagacacaggactacccatcagtgccca cgctggttcgcctgcttgccaagcataacatcatccccatctttgctgtcaccaactactcttacagctactatgagaagc tccataagtatttccccgtctcctctctgggcgtcctgcaggaggattcatccaacatcgtggagctgctggaggaggcct tctatcgaattcgctccaacctggacatccgggctctggacagccccagaggcctgagaacagaggtcacctccgatactc tccagaagacggagactgggtcctttcacatcaagcgggggaagtgggcacatacaatgtgcatctccgggcagtggagg 45 acatagalgggacacatgtgtgccagctggctaaagaacaaggcggcaacatccacctgaaaccctccttctctgatg gcctccggatggacgcgagtgtgatctgtgacgtgtgccctgtgagctgcaaaaggaagttcgatcagctcgctgtcact t cagaggagact t catgtgtgtgcactgtgtgtgcaatgagggctggagtggcaaaacctgcaactgctccaccggctctctgagtgacacacagccetgcctgcgtgagggtgaggacaaaccgtgctcgggccacggcgagtgccagtgcggacgctgtg tgtgctatggtgaaggccgctacgagggtcacttctgcgagtatgacaacttccagtgtccccggacctctggattcctgt gcaatgccacctgcatcgatagcaacgggggcatctgcaacggccgaggctactgtgagtgtggccgttgtcactgcaacc agcagtcgctctacacggacaccacctgtgagatcaactactctgcgatactgggtctctgtgaggatctccgctcctgcg tacagtgccaggcctggggcaccggggagaagaaagggcgcgcgtgtgacgattgcccctttaaagtcaagatggtagacg agcttaagaaagaagaggtggtggagtactgctccttccgggatgaggatgacgactgcacttacagctacaacgtggagg gcctggggcttcttccttgctgcaaccgaggtcacatggtgggctttaaggaagatcactatatgcttcgggagaacctga tggcctctgaccacctggacacgcccatgctacgaagcgggaacctcaagggacgagacacagtccgctggaagatcacca acaatqtqcaqcqccctqqctttqccacccatqccqccaqcaccaqccccacqqaqctcqtaccctacqqqctqtccctqc 60 gccttggccgcctctgcactgagaaccttatgaagccgggcacccgagagtgtgaccagctacgccaggaggtggaggaaa atctgaatgaggtgtatagacaggtcagcggcgcacacaagctccagcaqacgaagttccgacagcagcccaacgccggga aaaagcaagaccacaccattgtggacacagtgttgctggcgcccgctccgccaagcagatgctgctgaagctgacagaga agcaggtggagcaggggtccttccatgaactgaaggtggcccctggctactacactgtcacggcagagcaggatgcccggg gcatggtggagttccaggagggcgtggagctggtggatgtgcgagtgcccctcttcatccggcctgaggatgatgatgaga 65 agcagetgetggtggaggecattgatgteectgtgageactgecacecttggtegeegtetggtaaacateaccattatea aggaacaagctagtggggtagtgtccttcgagcagcctgaatactcggtgagtcgtggagaccaggtggcccgcatccctg

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8. SEQ ID NO:8 Murine integrin beta4 protein sequence:

- MAGPCCSPWV KLLLLARMLS ASLPGDLANR CKKAQVKSCT ECIRVDKSCA YCTDELFKERRCNTQADVLA 35 AGCRGESILV MESSLEITEN TQIVTSLHRS QVSPQGLQVR LRRGEERTFVFQVFEPLESP VDLYILMDFS NSMSDDLDNL KQMGQNLAKI LRQLTSDYTI GFGKFVDKVSVPQTDMRPEK LKEPWPNSDP PFSFKNVISL TENVEEFWNK LQGERISGNL DAPEGGFDAILQTAVCTRDI GWRADSTHLL VFSTESAFHY EADGANVLAG IMNRNDEKCH LDASGAYTQY KTQDYPSVPT LVRLLAKHNI IPIFAVTNYS YSYYEKLHKY FPVSSLGVLQ EDSSNIVELLEEAFYRIRSN 40 LDIRALDSPR GLRTEVTSDT LQKTETGSFH IKRGEVGTYN VHLRAVEDIDGTHVCQLAKE DQGGNIHLKP SFSDGLRMDA SVICDVCPCE LQKEVRSARC HFRGDFMCGHCVCNEGWSGK TCNCSTGSLS DTQPCLREGE DKPCSGHGEC QCGRCVCYGE GRYEGHFCEYDNFQCPRTSG FLCNDRGRCS MGECVCEPGW TGRSCDCPLS NATCIDSNGG ICNGRGYCEC GRCHCNQQSL YTDTTCEINY SAILGLCEDL RSCVQCQAWG TGEKKGRACD DCPFKVKMVDELKKEEVVEY 45 CSFRDEDDDC TYSYNVEGDG SPGPNSTVLV HKKKDCLPAP SWWLIPLLIFLLLLLLLLL LCWKYCACCK ACLGLLPCCN RGHMVGFKED HYMLRENLMA SDHLDTPMLRSGNLKGRDTV RWKITNNVQR PGFATHAAST SPTELVPYGL SLRLGRLCTE NLMKPGTRECDQLRQEVEEN LNEVYRQVSG AHKLQQTKFR QQPNAGKKQD HTIVDTVLLA PRSAKQMLLK LTEKQVEQGS FHELKVAPGY YTVTAEQDAR GMVEFQEGVE LVDVRVPLFI RPEDDDEKQLLVEAIDVPVS 50 TATLGRRLVN ITIIKEQASG VVSFEQPEYS VSRGDQVARI PVIRHILDNGKSQVSYSTQD NTAHGHRDYV PVEGELLFHP GETWKELQVK LLELQEVDSL LRGRQVRRFQVQLSNPKFGA RLGQPSTTTV ILDETDRSLI NQTLSSPPPP HGDLGAPQNP NAKAAGSRKIHFNWLPPPGK PMGYRVKYWI QGDSESEAHL LDSKVPSVEL TNLYPYCDYE MKVCAYGAKG EGPYSSLVSC RTHQEVPSEP GRLAFNVVSS TVTQLSWAEP AETNGEITAY EVCYGLVNEDNRPIGPMKKV
 - SSTLTRDYHS LTRTEHSHSG TLPRDYSTLTSLSSQASLLS GKMGGAGFRC PGLLGP

 9. SEQ 1D NO:9 Mouse beta4 dominant negative cds:

55 LVDNPKNRML LIENLRDSQP YRYTVKARNG AGWGPEREAI INLATQPKRPMSIPIIPDIP IVDAQGGEDY ENFLMYSDDV LRSPASSQRP SVSDDTEHLV NGRMDFAYPGSANSLHRMTA ANVAYGTHLS PHLSHRVLST

aaactgaaggagcctggcccaacagtgatcccccgttctccttcaagaacgttatcagcttaacggagaatgtggaagaa ttctqqaacaaactgcaaggagaacgcatctcaggcaacctggacgctcctgaagggggctttgatgccatcctgcagaca gaggctgatggtgccaacgttctggccggcatcatgaaccgcaatgatgagaaatgccacctggacgcctcgggcgcctac 5 acccaatacaagacacaggactacccatcagtgcccacgctggttcgcctgcttgccaagcataacatcatcaccatcttt qctqtcaccaactactcttacagctactatgagaagctccataagtatttccccgtctcctctctgggcgtcctgcaggag qattcatccaacatcgtggagctgctggaggaggccttctatcgaattcgctccaacctggacatccgggctctggacagc cccagaggcctgagaacagaggtcacctccgatactctccagaagacggagactgggtcctttcacatcaagcgggggsaa gtgggcacatacaatgtgcatctccgggcagtggaggacatagatgggacacatgtgtgccagctggctaaagaagaccaa 10 gggggcaacatccacctgaaaccctccttctctgatggcctccggatggacgcgagtgtgatctgtgacgtgtgcccctgt gacaacttecagtgteceeggacetetggattectgtgeaatgaceggggaegetgttetatgggagagtgtgtgtgtgtgag 15 cctggttggacaggccgcagctgcgactgtcccctcagcaatgccacctgcatcgatagcaacgggggcatctgcaacggc cgaggctactgtgagtgtgggccgttgtcactgcaaccagcagtcgctctacacggacaccacctgtgagatcaactactct qcqatactqqqtctctqtqaggatctccgctcctgcgtacagtgccaggcctggggcaccggggagaagaaagggcgcgcg tgtgacgattgcccctttaaagtcaagatggtagacgagcttaagaagaagaggtggtggagtactgctccttccgggat gaggatgacgactgcacttacagctacaacgtggagggcgacggcagccctgggcccaacagcacagtcctggtccacaaa 20 aagaaagactgcctcccggctccttcctggtggctcatccccctgctcatcttcctcctgttgctcctggcgttgcttctg ctgctctgctggaaatga

10. SEQ ID NO:10 Mouse beta4 dominant negative protein sequence:

MAGPCCSPWV KLLLARMLS ASLPGDLANR CKKAQVKSCT ECIRVDKSCA YCTDELFKERRCNTQADVLA
AGCRGESILV MESSLEITEN TQIVTSLHRS QVSPQGLQVR LRRGEERTFVFQVFEPLESP VDLYILMDFS

NSMSDDLDNL KQMGQNLAKI LRQLTSDYTI GFGKFVDKVSVPQTDMRPEK LKEPWPNSDP PFSFKNVISL
TENVEEFWNK LQGERISGNL DAPEGGFDAILQTAVCTRDI GWRADSTHLL VFSTESAFHY EADGANVLAG
IMNRNDEKCH LDASGAYTQY
KTQDYPSVPT LVRLLAKHNI IPIFAVTNYS YSYYEKLHKY FPVSSLGVLQ EDSSNIVELLEEAFYRIRSN
LDIRALDSPR GLRTEVTSDT LQKTETGSFH IKRGEVGTYN VHLRAVEDIDGTHVCQLAKE DQGGNIHLKP
SFSDGLRMDA SVICDVCPCE LQKEVRSARC HFRGDFMCGHCVCNEGWSGK TCNCSTGSLS DTQPCLREGEDKPCSGHGEC QCGRCVCYGE GRYEGHFCEYDNFQCPRTSG FLCNDRGRCS MGECVCEPGW TGRSCDCPLS
NATCIDSNGG ICNGRGYCEC
GRCHCNQQSL YTDTTCEINY SALLGLCEDL RSCVQCQAWG TGEKKGRACD DCPFKVKMVDELKKEEVVEY
CSFRDEDDDC TYSYNVEGDG SPGPNSTVLV HKKKDCLPAP SWWLIPLLIFLLLLLALLLLLLCWK

35

11. SEQ ID NO:11 Murine laminin gamma2 chain complete cds (acc# U43327):

40 cagcatcagagcaagagccgctgcctaccctgcaactgccactcaaagggttccctcagtgctggatgtgaca actotggacaatgcaggtgtaagccaggtgtgacaggacaaagatgtgaccagtgtcagccaggcttccatat gctcaccgatgctggatgcacccgagaccaggggcaactagattccaagtgtgactgtgacccagctggcatc tctggaccetgtgattctggccgatgtgtctgcaaaccagccgtcactggagagcgctgtgataggtgccgac cacqtqactatcatctqqaccqqqcaaaccctqagggctgtacccagtgtttctgctatgggcattcagccag gcggttcagagaaacggggcacctgcaaaactccactggtcacagcgccatcgggacgtgtttagttctgccc gaagatcagaccccgtctatttcgtggcccctgccaaattcctcggtaaccagcaagtgagttacgggcagag cctgtcttttgactaccgcgtggacagaggaggtagacagccgtctgcctacgatgtgatcctggaaggtgct ggtctacagatcagagctcctctgatggctccaggcaagacacttccttgtgggatcacaaagacttacacat 50 tcaqactgaatqaacatccaagcagtcactggagtccccagctgagttatttcgaatatcgaaggttactgcg qaacctcacagccctcctqatqatccgagctacgtacggagaatatagtacagggtacattgataacgtgacc ctggtttcagcccgccctgtccttggagccccagccccttgggttgaacgttgtgtatgcctgctggggtaca agggacaattetgecaggaatgtgettetggttacaaaagagatteggeaagattgggegetttttggegeetg tgttccctgtaactgccaaggggaggggcctgtgatccagacacgggagattgctactcgggggacgagaat 55 cctgacattgagtgtgctgactgtcccatcggtttctacaatgacccacatgacccccgcagctgcaagccat gtccctgtcacaatgggttcagctgttcagtgatgcctgagacagaggaggtggtgtgtaacaactgtccccc tggggtcacaggtgcccgctgtgagctctgtgctgatggcttcttttggggatccctttggggaacatggccca gtgaggccttgtcaacgctgccaatgcaacaacaacgtggaccccaatgcctctgggaactgtgaccagttga caqqcaqatqcttqaaatqtatctacaacacqqccqqtqtctactgtgaccagtgcaaagcaggttactttgg 60 agacccattggctcccaacccagcagacaagtgtcgagcttgcaactgcagccccatgggtgcggagcctgga gagtgtcgaggtgatggcagctgtgtttgcaagccaggctttggcgccttcaactgtgatcacgcagccctaa ccagttgtcctgcttgctacaatcaagtgaagattcagatggaccagtttacccagcagctccagagcctgga

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12. SEQ ID NO:12 Murine laminin gamma2 chain protein sequence (acc# AAA85256)

1 mpalwisccl gvallipasq atsrrevcdc ngksrqcvfd qelhrqagsg frcincndnt 61 agvhcersre gfyqhqsksr clpcnchskg slsagcdnsg qcrckpgvtg qrcdqcqpgf 121 hmltdagetr dqgqldskcd cdpagisgpc dsgrcvckpa vtgercdrcr prdyhldran 30 181 pegctqcfcy ghsaschasa dfsvhkitst fsqdvdgwka vqrngapakl hwsqrhrdvf 241 ssarrsdpvy fvapakflgn qqvsygqsls fdyrvdrggr qpsaydvile gaglqirapl 301 mapgktlpcg itktytfrln ehpsshwspq lsyfeyrrll rnltallmir atygeystgy 361 idnytlysar pylgapapwy ercycllgyk gqfcqecasg ykrdsarlga fgacypcncq 421 gegacdpdtg dcysgdenpd iecadcpigf yndphdprsc kpcpchngfs csympeteev 35 481 vcnncppgvt garcelcadg ffgdpfgehg pvrpcqrcqc nnnvdpnasg ncdqltgrcl 541 kciyntagvy cdqckagyfg dplapnpadk cracncspmg aepgecrgdg scvckpgfga 601 fncdhaalts cpacynqvki qmdqftqqlq slealvskaq ggggggtvpv qlegrieqae 661 qalqdilgea qisegamrav avrlakarsq endyktrldd lkmtaerira lgsqhqnrvq 721 dtsrlisqmr lslagseall entnihsseh yvgpndfksl aqeatrkads haesanamkq 40 781 laretedysk qalslarkl1 sggggsgswd ssvvgglmgk lektkslsqq lslegtqadi 841 eadrsyghsl rlldsasglq gvsdlsfqve akrirqkads lsnlvtrqtd aftrvrnnlg 901 nweketrqll qtgkdrrqts dqllsranla knraqealsm gnatfyeven ilknlrefdl 961 qvedrkaeae eamkrissis qkvadasdkt qqaetalgsa tadtqrakna arealeisse 1021 ieleigsln1 eanvtadgal amekgtatlk semremiela rkelefdtdk dtvqlvitea 45 1081 gqadaratsa gvtiqdtlnt 1dgilhlidq pgsvdeegmm lleqglfqak tqinsrlrpl 1141 msdleervrr qrnhlhllet sidgiladvk nlenirdnlp pgcyntqale qq

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13. SEQ ID NO:13 Human laminin gamma2 chain complete cds (acc# AH006634)

gttgaacagtgtatatgtcctgttgggtacaaggggcaattctgccaggattgtgcttctggctacaagagagattcagcg agactggggccttttgggcacctgtattccttgtaactgtcaagggggaggggctgtgatccagacacaggagattgttat tcaqqqqatqaqaatcctqacattqaqtqtqctqactgcccaattggtttctacaacgatccgcacgacccccgcagctgc aagccatgtcctgtcataacgggttcagctgctcagtgatgccggagacggaggtggtgttgcaataactgcctccc 5 ggggtcaccggtgcccgctgtgagctctgtgctgatggctactttggggacccctttggtgaacatggcccagtgaggcct tgtatccacaacacagccggcatctactgcgaccagtgcaaagcaggctacttcggggacccattggctcccaacccagca gacaagtgtcgagcttgcaactgtaaccccatgggctcagagcctgtaggatgtcgaagtgatggcacctgtgtttgcaag ccaggatttggtggccccaactgtgagcattgagcattcagctgtccagcttgctataatcaagtgaagattcagatggat 10 cagtttatgcagcagcttcagagaatggaggcctgatttcaaaggctcagggtggtgatggagtagtacctgatacagag ctggaaggcaggatgcagccggctgagcaggcccttcaggacattctgagagatgcccagatttcagaaggtgctagcaga tecettggtetceagttggceaaggtgaggagceaagagaacagetaceagageegcettggatgaceteaagatgactgtg gaaagagttcgggctctgggaagtcagtaccagaaccgagttcgggatactcacaggctcatcactcagatgcagctgagc ctggcagaaagtgaagcttccttgggaaacactaacattcctgcctcagaccactacgtggggccaaatggctttaaaagt 15 ctqqctcagqaggccacaagattagcagaaagccacgttgagtcagccagtaacatggagcaactgacaagggaaactgag gactattccaaacaagccctctcactggtgcgcaaggccctgcatgaaggagtcggaagcggaagcggtagcccggacggt gctgtggtgcaagggcttgtggaaaaattggagaaaaccaagtccctggcccagcagttgacaagggaggccactcaagcg gaaattgaagcagataggtcttatcagcacagtctccgcctcctggattcagtgtctcggcttcagggagtcagtgatcag teettteaygtggaagaageaaagaggateaaacaaaaageggatteaeteteaaegetggtaaccaggeatatggatgag 20 ttcaagcgtacacaaaagaatctgggaaactggaaagaagaagcacagcactcttacagaatggaaaaagtgggagagag aaatcagatcagctgctttcccgtgccaatcttgctaaaagcagagcacaagaagcactgagtatgggcaatgccactttt tatqaaqttqaqaqcatccttaaaaacctcaqagaqtttgacctqcaggtggacaacagaaaagcagaagctgaagaagcc gctgctgctgatgcacagagggcaaagaatggggccggggaggccctggaaatctccagtgagattgaacaggagattggg 25 agtctgaacttggaagccaatgtgacagcagatggagccttggccatggaaaagggactggcctctctgaagagtgagatg agggaagtggaaggagctggaaaggaaggagctggagtttgacacgaatatggatgcagtacagatggtgattacagaa gcccagaaggttgataccagagccaagaacgctggggttacaatccaagacacactcaacacttagacggcctcctgcat ctgatggaccagcctctcagtgtagatgaagaggggctggtcttactggagcagaagctttcccgagccaagacccagatc aacagccaactgcggcccatgatgtcagagctggaagagagggcacgtcagcagaggggccacctccatttgctggagaca 30 agcatagatqqqattctqqctqatqtqaaqaacttqqaqaacattaqqqacaacctqcccccaqqctqctacaatacccaq gctcttgagcaacagtga

14. SEQ ID NO:14 Human laminin gamma2 chain protein sequence (acc# AAC50457) Alternative splice form

```
1 mpalwlgccl cfslllpaar atsrrevcdc ngksrqcifd relhrqtgng frclncndnt
35 61 dgihcekckn gfyrhrerdr clpcncnskg slsarcdnsg resckpgvtg arcdrelpgf
    121 hmltdagctq dqrlldskcd cdpagiagpc dagrcvckpa vtgercdrcr sgyynldggn
   181 peqctqcfcy qhsascrssa eysvhkitst fhqdvdgwka vqrngspakl qwsqrhqdvf
   241 ssaqrldpvy fvapakflgn qqvsygqsls fdyrvdrggr hpsahdvile gaglritapl
301 mplgktlpcg ltktytfrln ehpsnnwspq lsyfeyrrll rnltalrira tygeystgyi
40 351 dnvtlisarp vsgapapwve qcicpvgykg qfcqdcasgy krdsarlgpf gtcipcncqg
    421 ggacdpdtgd cysgdenpdi ecadcpigfy ndphdprsck pcpchngfsc sympeteevv
    481 cnncppgvtg arcelcadgy fgdpfgehgp vrpcqpcqcn nnvdpsasgn cdrltgrclk
   541 cihntagiyc dqckagyfgd plapnpadkc racncnpmgs epvgcradgt cvckpgfggp
    601 ncehgafscp acynqvkiqm dqfmqqlqrm ealiskaggg dgvvpdtele grmqqaeqal
45 661 qdilrdaqis egasrslglq lakvrsqens yqsrlddlkm tvervralgs qyqnrvrdth
    721 rlitqmqlsl aeseaslgnt nipasdhyvg pngfkslaqe atrlaeshve sasnmeqltr
   781 etedyskaal slvrkalheg vgsgsgspdg avvaglvekl ektkslagal treatgaeie
   841 adrsyghslr lldsvsrlgg vsdqsfqvee akrikqkads lstlvtrhmd efkrtqknlg
901 nwkeeaqqll qngksgreks dqllsranla ksraqealsm gnatfyeves ilknlrefdl
50 961 qvdnrkaeae eamkrlsyis qkvsdasdkt qqaeralgsa aadaqrakng agealeisse
    1021 iegeigslnl eanvtadgal amekglaslk semrevegel erkelefdtn mdavqmvite
   1081 agkvdtrakn agvtigdtln tldgllhlmd qplsvdeegl vllegklsra ktqinsqlrp
   1141 mmseleerar qqrghlhlle tsidgiladv knlenirdnl ppgcyntqal eqq
```

15. SEQ ID NO:15 Human laminin gamma2 chain protein sequence (acc# AAC50456) Alternative splice form

1 mpalwlgccl cfslllpaar atsrrevede ngksrqcifd relhrqtgng frelnendnt
61 dgiheeken gfyrhrerdr elpenenskg slsarednsg resekpgvtg aredrelpgf
121 hmltdagetq dqrlldsked edpagiagpe dagrevekpa vtgeredrer sgyynldggn
181 pegetqefey ghsaserssa eysvhkitst fhqdvdgwka vqrngspakl qwsqrhqdvf
60 241 ssaqrldpvy fvapakflgn qqvsygqsls fdyrvdrggr hpsahdvile gaglritapl
301 mplgktlpcg ltktytfrln ehpsnnwspq lsyfeyrrll rnltalrira tygeystgyi
361 dnvtlisarp vsgapapwve qeiepvgykg qfeqdeasgy krdsarlgpf gteipenegg
421 ggaedpdtgd cysgdenpdi ecadepigfy ndphdprsek pepchngfse svmpeteevv
481 enneppgvtg arceleadgy fgdpfgehgp vrpcqpcqen nnvdpsasgn edrltgrelk
65 541 eihntagiye dqekagyfgd plapnpadke racnenpmgs epvgersdgt evekpgfggp

55

601 ncehgafscp acynqvkiqm dqfmqqlqrm ealiskaqgg dgvvpdtele grmqqaeqal
661 qdilrdaqis egasrslglq lakvrsqens yqsrlddlkm tvervralgs qyqnrvrdth
721 rlitqmqlsl aeseaslgnt nipasdhyvg pngfkslaqe atrlaeshve sasnmeqltr
781 etedyskqal slvrkalheg vgsgsgspdg avvgglvekl ektkslaqql treatqaeie
841 adrsyqhslr lldsvsrlgg vsdqsfqvee akrikqkads lstlvtrhmd efkrtqknlg
901 nwkeeaqqll qngksgreks dqllsranla ksraqealsm gnatfyeves ilknlrefdl
961 qvdnrkaeae eamkrlsyis qkvsdasdkt qqaeralgsa aadaqrakng agealeisse
1021 ieqeigslnl eanvtadgal amekglaslk semrevegel erkelefdtn mdavqmvite
1081 aqkvdtrakn agvtiqdtln tldgllhlmg m

16. Degenerate Human integrin beta4 subunit cds

10

atg gca gCg cca cgc ccc agc cca tgg gcc agg ctg ctc ctg gca gcc ttg atc agc gtc ago cto tot ggg aco ttg aac cgc tgc aag aag gcc cca gtg aag ago tgc acg gag tgt gtc cgt gtg gat aag gac tgc gcc tac tgc aca gac gag atg ttc agg gaccggcgctgcaacacccaggcggagctgctggccgcggggctgccagcgggagagcatcgtggtcatggagagcagcttc 15 caaatcacagaggagacccagattgacaccaccctgcggcgcagccagatgtccccccaaggcctgcgggtccgtctgcgg cccggtgaggagcggcattttgagctggaggtgtttgagccactggagagccccgtggacctgtacatcctcatggacttc tccaactccatgtccgatgatctggacaacctcaagaagatggggcagaacctggctcgggtcctgagccagctcaccagc gactacactattggatttggcaagtttgtggacaaagtcagcgtcccgcagacggacatgaggcctgagaagctgaaggag ccttggcccaacagtgaccccccttctccttcaagaacgtcatcagcctgacagaagatgtggatgagttccggaataaa 20 ctgcagggagagcggatctcaggcaacctggatgctcctgagggcggcttcgatgccatcctgcagacagctgtgtgcacg agggacattggctggcgcccggacagcacccacctgctggtcttctccaccgagtcagccttccactatgaggctgatggc gccaacgtgctggctggcatcatgagccgcaacgatgaacggtgccacctggacaccacgggcacctacacccagtacagg acacaggactaccegtcggtgcccaccctggtgcgcctgctcgccaagcacaacatcatccccatctttgctgtcaccaac tactcctatagctactacgagaagcttcacacctatttccctgtctcctcactgggggtgctgcaggaggactcgtccaac 25 atcgtggagctgctggaggaggccttcaatcggatccgctccaacctggacatccgggccctagacagccccgaggcctt cggacagaggtcacctccaagatgttccagaagacgaggactgggtcctttcacatccggcgggggaagtgggtatatac caggtgcagctgcgggcccttgagcacgtggatgggacgcacgtgtgccagctgccggaggaccagaagggcaacatccat ctgaaaccttccttctccgacggcctcaagatggacgcgggcatcatctgtgatgtgtgcacctgcgagctgcaaaaagag gagtgccagtgcgggcactgtgtgtgctacggcgaaggccgctacgagggtcagttctgcgagtatgacaacttccagtgt ccaagctgtgactgtcccctcagcaatgccacctgcatcgacagcaatgggggcatctgtaatggacgtggccactgtgag tgtggccgctgccactgccaccagcagtcgctctacacggacaccatctgcgagatcaactactcggcgtccacccgggcc 35 tctgcgaggacctacgctcctgcgtgcagtgccaggcgtggggcaccggcgagaagaagggggcgcacgtgtgaggaatgca acttcaaggtcaagatggtggacgagcttaagagaggcgaggaggtggtggtggtgcctccttccgggacgaggatgacg actgcacctacagctacaccatggaaggtgacggcgcccttgggcccaacagcactgtcctggtgcacaagaagaagggac tgccctccgggctccttctggtggctcatccccctgctcctcctcctcctgctcctggccctgctactgctactgctatgc tggaagtactgtgcctgctgcaaggcctgcctggcacttctcccgtgctgcaaccgaggtcacatggtgggctttaaggaa 40 gaccactacatgctgegggagaacctgatggcctctgaccacttggacacgcccatgctgcgcagcgggaacctcaagggc cgtgacgtggtccgctggaaggtcaccaacaacatgcagcggcctggctttgccactcatgccgccagcatcaaccccaca gagctggtgccctacgggctgtccttgcgcctggcccgcctttgcaccgagaacctgctgaagcctgacactcgggagtgc gcccagctgcgccaggaggtggaggagaacctgaacgaggtctacaggcagatctccggtgtacacaagctccagcagacc aagttccggcagcagcccaatgccgggaaaaagcaagaccacaccattgtggacacagtgctgatggcgccccgctcggcc 45 aagccggccctgctgaagcttacagagaagcaggtggaacagagggccttccacgacctcaaggtggcccccggctactac acceteactgcagaccaggacgcccggggcatggtggagttccaggagggcgtggagctggtggacgtacgggtgcccctc tttatccggcctgaggatgacgagaagcagctgctggtggaggccatcgacgtgcccgcaggcactgccaccctcggc cgccgcctggtaaacatcaccatcatcaaggagcaagccagagacgtggtgtcctttgagcagcctgagttctcggtcagc cgcggggaccaggtggcccgcatccctgtcatccggcgtgtcctggacggcgggaagtcccaggtctcctaccgcacacag 50 gatggcaccgcgcagggcaaccgggactacatccccgtggagggtgagctgctgttccagcctgggaggcctggaaagag agcaaccctaagtttggggccacctgggccagcccactccaccaccatcatcatcagggacccagatgaactggaccgg agetteacgagteagatgttgteateacagecacccctcacggcgacctgggcgccccgcagaaccccaatgctaaggcc gctgggtccaggaagatccatttcaactggctgcccccttctggcaagccaatggggtacagggtaaagtactggattcag 55 ggtgactccgaatccgaagcccacctgctcgacagcaaggtgccctcagtggagctcaccaacctgtacccgtattgcgac gtgcccagcgagccagggcgtctggccttcaatgtcgtctcctccacggtgacccagctgagctgagctgagccggctgag accaacggtgagatcacagcctacgaggtctgctatggcctggtcaacgatgacaaccgacctattgggcccatgaagaaa gtgctggttgacaaccctaagaaccggatgctgcttattgagaaccttcgggagtcccagccctaccgctacacggtgaag 60 gcgcgcaacggggccggctgggggcctgagcgggaggccatcatcaacctggccacccagcccaagaggcccatgtccatc cccatcatccctgacatccctatcgtggacgcccagagcggggaggactacgacagcttccttatgtacagcgatgacgtt ctacgctctccatcgggcagccagaggcccagcgtctccgatgacactggctgcggctggaagttcgagccctgctgggg gaggagetggacetgeggegegteacgtggeggetgececeggageteatecegegeetgteggeeageageggegetee 65 cccgggccccccggagagcacctggtgaatggccggatggactttgccttcccgggcagcaccaactccctgcacaggatg accacgaccagtgctgctgcctatggcacccacctgagcccacacgtgccccaccgcgtgctaagcacatcctccaccctc

17. SEQ ID NO:17 Human beta4 integrin protein sequence Variant V at 34 to I:

1 magprpspwa rlllaalisv slsgtlnrck kapIksctec vrvdkdcayc tdemfrdrrc 61 ntqaellaag cqresivvme ssfqiteetq idttlrrsqm spqglrvrlr pgeerhfele 20 121 vfeplespvd lyilmdfsns msddldnlkk mgqnlarvls qltsdytigf gkfvdkvsvp 181 qtdmrpeklk epwpnsdppf sfknvislte dvdefrnklg gerisgnlda peggfdailq 241 tavctrdigw rpdsthllvf stesafhyea dganvlagim srnderchld ttgtytqyrt 301 qdypsvptlv rllakhniip ifavtnysys yyeklhtyfp vsslgvlqed ssnivellee 361 afnrirshld iraldsprgl rtevtskmfq ktrtgsfhir rgevgiyqvq lralehvdgt 25 421 hvcqlpedqk gnihlkpsfs dglkmdagii cdvctcelqk evrsarcsfn gdfvcgqcvc 481 segwsgqtcn cstgsladiq pclregedkp csgrgecqcg hcvcygegry egqfceydnf 541 qcprtsgflc ndrgrcsmgq cvcepgwtgp scdcplsnat cidsnggicn grghcecgrc 601 hchqqslytd ticeinysas trasartyap acsarrqapa rrrqarvrna tsrsrwwts1 661 rearrwwcaa psgtrmttap tatpwkvtap lgptalswct rrrdcppgsf wwliplllll 30 721 lpllallll1 cwkycaccka clallpccnr ghmvgfkedh ymlrenlmas dhldtpmlrs 781 gnlkgrdvvr wkvtnnmqrp gfathaasin ptelvpygls lrlarlcten llkpdtreca 841 qlrqeveenl nevyrqisgv hklqqtkfrq qpnagkkqdh tivdtvlmap rsakpallkl 901 tekqveqraf hdlkvapgyy tltadqdarg mvefqegvel vdvrvplfir pedddekqll 961 veaidvpagt atlgrrlvni tiikeqardv vsfeqpefsv srgdqvarip virrvldggk 35 1021 sqvsyrtqdg taggnrdyip vegellfqpg eawkelqvkl lelqevdsll rgrqvrrfhv 1081 qlsnpkfgah lgqphsttii irdpdeldrs ftsqmlssqp pphqdlqapq npnakaaqsr 1141 kihfnwlpps gkpmgyrvky wiqgdsesea hlldskvpsv eltnlypycd yemkvcayga 1201 ggegpysslv scrthqevps epgrlafnvv sstvtqlswa epaetngeit ayevcyglvn 1261 ddnrpigpmk kvlvdnpknr mllienlres qpyrytvkar ngagwgpere aiinlatqpk 40 1321 rpmsipiipd ipivdaqsge dydsflmysd dvlrspsgsq rpsvsddtgc gwkfepllge 1381 eldlrrvtwr lppeliprls assgrssdae aphgppddgg aggkggslpr satpgppgeh 1441 lvngrmdfaf pgstnslhrm tttsaaaygt hlsphvphrv lstsstltrd ynsltrsehs 1501 hsttlprdys tltsvsshgl ppiwehgrsr lplswalgsr sraqmkgfpp srgprdsiil 1561 agrpaapswg pdsrltagvp dtptrlvfsa lgptslrvsw qeprcerplq gysveyqlln 45 1621 ggelhrlnip npaqtsvvve dllpnhsyvf rvraqsqegw greregviti esqvhpqspl 1681 cplpgsaftl stpsapgplv ftalspcslq lswerprrpn gdivgylvtc emaggggpat 1741 afrvdgdspe srltvpglse nvpykfkvqa rttegfgper egiitiesqd ggpfpqlgsr 1801 aglfqhplqs eyssisttht satepflvgp tlgaqhleag gsltrhvtqe fvsrtlttsg 1861 tlsthmdqqf fqt

18. SEQ ID NO:18 Human integrin beta4 subunit cds Variant at 34 V to I:

cggacagaggtcacctccaagatgttccagaagacgaggactgggtcctttcacatccggcggggggaagtgggtatatac caggtgcagctgcgggcccttgagcacgtggatgggacgcacgtgtgccagctgccggaggaccagaagggcaacatccat ctgaaaccttccttctccgacggcctcaagatggacgcgggcatcatctgtgatgtgtgcacctgcgagctgcaaaaagag gagtgccagtgcgggcactgtgtgtgctacggcgaaggccgctacgagggtcagttctgcgagtatgacaacttccagtgt ccaagctgtgactgtcccctcagcaatgccacctgcatcgacagcaatgggggcatctgtaatggacgtggccactgtgag tgtggccgctgccactgccaccagcagtcgctctacacggacaccatctgcgagatcaactactcggcgtccacccgggcc 10 tetgegaggacetaegeteetgegtgeagtgeeaggegtggggeaceggegagaagaaggggegeaegtgtgaggaatgea acttcaaggtcaagatggtggacgagcttaagagaggcgaggaggtggtggtggtgctgctccttccgggacgaggatgacg actgcacctacagctacaccatggaaggtgacggcgccctgggcccaacagcactgtcctggtgcacaagaagaagggac tgccctccgggctccttctggtggctcatccccctgctcctcctcctcctgctcctggccctgctactgctactgctatgc tggaagtactgtgcctgctgcaaggcctgcctggcacttctcccgtgctgcaaccgaggtcacatggtgggctttaaggaa 15 gaccactacatgctgcgggagaacctgatggcctctgaccacttggacacgcccatgctgcgcagcgggaacctcaagggc cgtgacgtggtccgctggaaggtcaccaacaacatgcagcggcctggctttgccactcatgccgccagcatcaaccccaca gagetggtgccctacgggctgtccttgcgcctggcccgcctttgcaccgagaacctgctgaagcctgacactcgggagtgc gcccagctgcgccaggaggtggaggagaacctgaacgaggtctacaggcagatctccggtgtacacaagctccagcagacc aagttccggcagcagcccaatgccgggaaaaagcaagaccacaccattgtggacacagtgctgatggcgccccgctcggcc 20 aagccggccttgctgaagcttacagagaagcaggttggaacagaggccttccacgacctcaaggttggcccccggctactac accetcaetgeagaccaggacgeecggggeatggtggagttecaggagggegtggagetggtggaegtaegggtgeecete tttatccggcctgaggatgacgacgagaagcagctgctggtggaggccatcgacgtgcccgcaggcactgccaccctcggc cgccgcctggtaaacatcaccatcatcaaggagcaagccagagacgtggtgtcctttgagcagcctgagttctcggtcagc cgcggggaccaggtggcccgcatccctgtcatccggcgtgtcctggacggcgggaagtcccaggtctcctaccgcacacag 25 gatggcaccgcgcagggcaaccgggactacatccccgtggagggtgagctgctgttccagcctgggagggcctggaaagag agcaaccctaagtttggggcccacctgggccagcccactccaccaccatcatcatcagggacccagatgaactggaccgg agcttcacgagtcagatgttgtcatcacagccacccctcacggcgacctgggcgccccgcagaaccccaatgctaaggcc gctgggtccaggaagatccatttcaactggctgccccttctggcaagccaatggggtacagggtaaagtactggattcag 30 ggtgactccgaatccgaagcccacctgctcgacagcaaggtgccctcagtggagctcaccaacctgtacccgtattgcgac głgcccagcgagccagggcgłcłggccłłcaałgłcgłctcccccacggłgacccagctgagcłgggcłgagccggcłgag accaacggtgagatcacagcctacgaggtctgctatggcctggtcaacgatgacaaccgacctattgggcccatgaagaaa gtgctggttgacaaccctaagaaccggatgctgcttattgagaaccttcgggagtcccagccctaccgctacacggtgaag 35 gcgcgcaacggggccggctgggggcctgagcgggaggccatcatcaacctggccacccagcccaagaggcccatgtccatc cccatcatccctgacatccctatcgtggacgcccagagcggggaggactacgacagcttccttatgtacagcgatgacgtt ctacgctctccatcgggcagccagaggcccagcgtctccgatgacactggctgcggctggaagttcgagcccctgctgggg gaggagctggacctgcggcgcgtcacgtggcggctgcccccggagctcatcccgcgcctgtcggccagcagcgggcgctcc 40 cccgggccccccggagagcacctggtgaatggccggatggactttgccttcccgggcagcaccaactccctgcacaggatg accacgaccagtgctgctgcctatggcacccacctgagcccacagtgcccaccgcgtgctaagcacatcctccaccctc acacgggactacaactcactgacccgctcagaacactcacactcgaccacactgccgagggactactccacctcacctccgtctcctcccacggcctccctcccatctgggaacacgggaggagcaggcttccgctgtcctgggccctggggtcccggagt 45 tggggcccagactctcgcctgactgctggtgtgcccgacacgcccacccgcctggtgttctctgccctggggcccacatct ctcagagtgagctggcaggagccgcggtgcgagcggccgctgcagggctacagtgtggagtaccagctgctgaacggcggt gagetgcateggetcaacatececaaccetgeccagaceteggtggtggtggaagaceteetgeecaaccactectaegtg ttccgcgtgcgggcccagagccaggaaggctggggccgagagcgtgagggtgtcatcaccattgaatcccaggtgcacccg cagagcccactgtgtcccctgccaggctccgccttcactttgagcactcccagtgccccaggcccgctggtgttcactgcc 50 ctgagcccagactcgctgcagctgagctgggagcggccacggaggcccaatggggatatcgtcggctacctggtgacctgt ctcagcgagaacgtgccctacaagttcaaggtgcaggccaggaccactgagggcttcggggccagagcgcgagggcatcatc accatagagtcccaggatggaggtcccttcccgcagctgggcagccgtgccgggctcttccagcacccgctgcaaagcgag tacagcagcatetecaccacccaccaccagcgccaccgagccettcctagtgggtccgaccetgggggcccagcacctggag 55 gcaggcggctccctcacccggcatgtgacccaggagtttgtgagccggacactgaccaccagcggaacccttagcacccac atggaccaacagttcttccaaacttga

19. SEQ ID NO:19 Degenerate Human integrin beta4 subunit cds variant at 34 Vto 1:

atg gca ggg cca cgG ccc agc cca tgg gcc agg ctg ctc ctg gca gcc ttg atc agc gtc

60 agc ctc tct ggg acc ttg aac cgc tgc aag aag gcc cca AtA aag agc tgc acg gag tgt

gtc cgt gtg gat aag gac tgc gcc tac tgc aca gac gag atg ttc agg

gaccggcgctgcaacaccccaggcggagctgctggcagcggggctgccagcgggagagcatcgtggtcatggagagcagcttc

caaatcacagaggagacccagattgacaccaccttgcggcgcagccagatgtccccccaaaggcctgcagggtccgtctgcagctgcagctgcagctggagagcctgtacatcctcatggacttc

65 tccaactccatgtccgatgatctggacaacctcaagaagatggggcagaacctggctcgggtcctgagacagctcaccagc

gactacactattggatttggcaagtttgtggacaaagtcagcgtcccgcagacggacatgaggcctgagaagctgaaagctgaagag

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What is claimed is:

1. A method of reducing the proliferation of a cancer cell which comprises inhibiting ligand binding to an integrin receptor on the cancer cell, wherein the integrin receptor comprises an integrin.

- 2. The method of claim 1, wherein the integrin receptor comprises integrin B4.
- 3. The method of claim 1, wherein the integrin receptor comprises integrin A6.
- 4. The method of claim 1, wherein the ligand that binds to the integrin receptor is laminin5.
 - 5. The method of claim 1, wherein the integrin receptor is A6B4.
 - 6. The method of claim 1, wherein the ligand comprises laminin5.
 - 7. The method of claim 1, wherein the ligand comprises the gamma-2 subunit of laminin5.
- 8. The method of claim 1, wherein inhibiting ligand binding to an integrin receptor comprises contacting a A6 integrin with a composition that inhibits ligand binding.
- 9. The method of claim 1, wherein inhibiting ligand binding to an integrin receptor comprises contacting a B4 integrin with a composition that inhibits ligand binding.
- 10. The method of claim 1, wherein inhibiting ligand binding to an integrin receptor comprises contacting a laminin5 with a composition that inhibits ligand binding.
- 11. The method of claim 1, wherein inhibiting ligand binding to an integrin receptor comprises contacting a gamma-2 subunit with a composition that inhibits ligand binding.
- 12. A method of reducing the proliferation of a cancer cell which comprises reducing integrin-integrin interaction, integrin receptor clustering interaction, or integrin-non-integrin protein interaction.
- 13. The method of claim 12, wherein reducing integrin-integrin interaction, integrin receptor clustering interaction, or integrin-non-integrin protein interaction comprises contacting a A6 integrin with a composition that inhibits an interaction between the B4 integrin and another integrin or protein molecule.
- 14. The method of claim 12, wherein reducing integrin-integrin interaction, integrin receptor clustering interaction, or integrin-non-integrin protein interaction comprises contacting a B4 integrin with a composition that inhibits the interaction between the B4 integrin and another integrin or protein molecule..
- 15. The method of claim 12, wherein reducing integrin-integrin interaction, integrin receptor clustering interaction, or integrin-non-integrin protein interaction comprises contacting a

laminin5 with a composition that inhibits ligand binding.

16. The method of claim 12, wherein reducing integrin-integrin interaction, integrin receptor clustering interaction, or integrin-non-integrin protein interaction comprises contacting a gamma2 subunit with a composition that inhibits ligand binding.

- 17. The method of claim 12, wherein the non-integrin protein comprises a growth factor receptor.
- 18. The method of claim 12, wherein the non-integrin protein comprises a hemidesomosome junction.
 - 19. The method of claim 12, wherein the non-integrin protein comprises a SH2 domain.
 - 20. The method of claim 19, wherein the non-integrin protein comprises a Shc protein.
 - 21. The method of claim 12, wherein the non-integrin protein comprises a IRS-1 protein.
 - 22. The method of claim 12, wherein the non-integrin protein comprises a IRS-2 protein.
- 23. A method of reducing the proliferation of a cancer cell which comprises reducing the production of an integrin by the cancer cell.
- 24. The method of claim 23, wherein the production of an integrin is reduced by inhibiting signaling leading to induction of expression of an integrin.
- 25. The method of claim 23, wherein reducing the production of an integrin comprises inhibiting alpha6 production.
- 26. The method of claim 25, wherein inhibiting alpha6 production further comprises using antisense molecules to alpha6 mRNA.
- 27. The method of claim 23, wherein reducing the production of an integrin comprises inhibiting beta4 production.
- 28. The method of claim 27, wherein inhibiting beta4 production further comprises using antisense molecules to beta4 mRNA.
- 29. A method of reducing the proliferation of a cancer cell which comprises reducing the production of an integrin receptor ligand by the cancer cell.
- 30. The method of claim 29, wherein reducing the production of an integrin receptor ligand comprises inhibiting gamma2 production.
- 31. The method of claim 29, wherein reducing the production of an integrin receptor ligand comprises inhibiting laminin production.
 - 32. The method of claim 29, wherein reducing the production of an integrin receptor

ligand comprises inhibiting laminin5 production.

33. A method of reducing the proliferation of a cancer cell which comprises interfering with an integrin signaling pathway.

- 34. The method of claim 33, wherein interfering with an integrin signaling pathway comprises contacting an A6 integrin in the cell with a composition that inhibits ligand binding.
- 35. The method of claim 33, wherein interfering with an integrin signaling pathway comprises contacting a B4 integrin with a composition that inhibits ligand binding.
- 36. The method of claim 33, wherein interfering with an integrin signaling pathway comprises contacting a laminin5 with a composition that inhibits ligand binding.
- 37. The method of claim 33, wherein interfering with an integrin signaling pathway comprises contacting a gamma2 subunit with a composition that inhibits ligand binding.
- 38. The method of claim 33, wherein interfering with an integrin signaling pathway comprises contacting the cancer cell with a molecule that interferes with at least one of talin, paxillin, vinculin, a CAS family protein, CRX, NCK, FAK, ILK, Src, Fyn, Shc, Grb-2, Guaning nucleotide exchange factors, SOS, DOCK 180, Vav, Syk, P-1-3 kinase, AKT, Bad, Bid, Caspase 9, Cdc42, PAK, Rac, Rho, Rho kinase, Ras, Caveolin, Tetraspan, Receptor-type protein tyrosine phosphatase, SHP-2, Alpha-actinin, Filamin, Cytohesin, Beta3-endonexin, ICAP-1, RACK-1, CIB, actin, receptor tyrosine kinase, IRS-1 or IRS-2.
- 39. The method of claim 33, wherein interfering with an integrin signaling pathway comprises contacting the cancer cell with an agent that interferes with post-translational modification of integrins.
- 40. The method of claim 39, wherein the post translational modification is glycosylation or phosphorylation.
 - 41. The method of claims 1, 12, 23, 29, 33, wherein the integrin comprises an A6 integrin.
 - 42. The method of claims 1, 12, 23, 29, 33, wherein the integrin comprises a B4 integrin.
- 43. The method of claims 1, 12, 23, 29, 33, further comprising reducing a laminin5-integrin interaction.
- 44. The method of claims 1, 12, 23, 29, 33, further comprising reducing a laminin5 gamma2 integrin interaction.
- 45. The method of claims 1, 12, 23, 29, 33, wherein reducing the proliferation of the cancer cells is selective.
 - 46. The method of claims 1, 12, 23, 29, 33, wherein the cancer cell is not an MDA-MB-

435 cell.

47. The method of claims 1, 12, 23, 29, 33, wherein the cancer cell is not an HMT-3522 cell.

- 48. The method of claims 1, 12, 23, 29, 33, wherein the cancer cell is not an RKO colon carcinoma line.
- 49. The method of claims 1, 12, 23, 29, 33, wherein the cancer cell does not express exogenous B4 integrin.
- 50. The method of claims 1, 12, 23, 29, 33, further comprising contacting the cancer cells with a small molecule, peptide, peptide mimetic, or oligonucleotide or synthetic analog thereof.
- 51. The method of claim 50, wherein the cancer cells are contacted with dominantnegative beta 4 integrin.
- 52. The method of claim 50, wherein the cancer cell is contacted with an antisense molecule.
- 53. The method of claim 52, wherein the antisense molecule is linked to a leader sequence which enables translocation across a cell membrane.
- 54. The method of claim 53, wherein the leader sequence binds to a cell surface protein which facilitates internalization.
- 55. The method of claim 54, wherein the leader sequence is TAT or antennapedia, or fragment thereof.
- 56. The method of claim 52, wherein the antisense molecule is an alpha6 RNA antisense molecule.
- 57. The method of claim 47, wherein the small molecule peptide, peptide mimetic, or oligonucleotide or synthetic analog thereof is linked to a carrier.
- 58. The method of claim 57, wherein the carrier is at least one of a lipidic carrier, charged carrier, retroviral carrier, TAT or fragment thereof, antennapedia or fragment thereof, or polyethylene glycol.
- 59. The method of any one of claims 1, 12, 23, 29, 33, further comprising contacting the cancer cell with another agent which modulates cell signaling, a chemotherapeutic drug, or treated with radiation or angiogenesis inhibitor.
- 60. The method of any of claims 1, 12, 23, 29, 33, wherein reducing the proliferation of cancer cell is *in vitro*.
 - 61. The method of any of claims 1, 12, 23, 29, 33, wherein reducing the proliferation

of the cancer cell is in vivo.

from the group consisting of melanoma, adenoma, lymphoma, myeloma, carcinoma, plasmocytoma, sarcoma, glioma, thyoma, leukemia, skin cancer, retinal cancer, breast cancer, prostate cancer, colon cancer, esophageal cancer, stomac cancer, pancreas cancer, brain tumors, lung cancer, ovarian cancer, cervical cancer, hepatic cancer, gastrointestinal cancer, and head and neck cancer cells.

- 63. The method of claims 1, 12, 23, 29, 33, wherein the cancer cell is killed.
- 64. The method of claims 1, 12, 23, 29, 33, wherein the cancer cell expresses a mutated Ras.
- 65. The method of claims 1, 12, 23, 29, 33, wherein the cancer cell expresses a mutated Ras and a mutated p53.
- 66. The method of claims 1, 12, 23, 29, 33, wherein the cancer cell expresses a mutated Ras and activates the AKT/PKB protein.
- 67. The method of claims 1, 12, 23, 29, 33, wherein the cancer cell expresses a mutated Ras, a mutated p53, and activates the AKT/PKB protein.
- 68. The method of claims 1, 12, 23, 29, 33, wherein the cancer cell expresses a mutated APC.
- 69. The method of claims 1, 12, 23, 29, 33, wherein the cancer cell expresses a mutated Ras and mutated APC.
- 70. A method of isolating molecules that bind with a target molecule selected from the group consisting, B4 integrin, alpha6 integrin, and the gamma2 subunit of laminin5 comprising 1) contacting a library of molecules with the target molecule and 2) collecting molecules that bind the target molecule producing an enriched population of molecules.
- 71. The method of claim 70, further comprising the step of repeating steps 1 and 2 with the enriched population of molecules.
- 72. The method of claim 70, wherein the library comprises a small molecule, peptide, peptide mimetic, or oligonucleotide.
- 73. A method of assessing a subject's risk of developing cancer comprising determining the amount of A6 present in a target cell obtained from the subject, wherein a determination of increased levels of A6 correlates with an increased risk of cancer.
 - 74. The method of claim 73, further comprising comparing the amount A6 present to the

amount in a control cell.

75. The method of claim 73, wherein determining the amount of A6 present in the target cell comprises assaying the amount of A6 mRNA in the cell.

- 76. The method of claim 75, wherein the assaying the amount of mRNA in the cell comprises hybridizing a A6 probe to a sample of the subject's mRNA.
- 77. The method claim 75, wherein the assaying the amount of mRNA in the cell comprises hybridizing a A6 primer to a sample of the subject's mRNA.
- 78. The method claim 77, wherein the assaying the amount of mRNA further comprises performing an nucleic acid amplification reaction involving the primer.
- 79. The method of claim 78, wherein the nucleic acid amplification reaction comprises reverse transcription, producing a cDNA.
- 80. The method of claim 79, wherein the nucleic acid amplification reaction further comprises performing a polymerase chain reaction on the cDNA.
- 81. The method of claim 73, wherein determining the amount of A6 present in the target cell comprises assaying the amount of A6 protein in the cell.
- 82. A method of assessing a subject's risk of acquiring cancer comprising determining the amount of B4 present in a target cell obtained from the subject, wherein a determination of increased levels of B4 correlates with an increased risk of cancer.
- 83. The method of claim 82, further comprising comparing the amount B4 present to the amount in a control cell.
- 84. A method of assessing a subject's risk of acquiring cancer comprising determining the amount of laminin5 present in a target cell obtained from the subject, wherein a determination of increased levels of laminin5 correlates with an increased risk of cancer.
- 85. The method of claim 84, further comprising comparing the amount laminin5 present to the amount in a control cell.
- 86. A method of assessing a subject's risk of acquiring cancer comprising determining the amount of gamma2 subunit present in a target cell obtained from the subject, wherein a determination of increased levels of gamma2 subunit correlates with an increased risk of cancer.
- 87. The method of claim 86, further comprising comparing the amount gamma2 subunit present to the amount in a control cell.
- 88. A method of reducing the proliferation of a cancer cell in a patient which comprises administering to the patient a composition which inhibits ligand binding to an integrin receptor on

the cancer cell, wherein the integrin receptor comprises an integrin.

- 89. A method of reducing the proliferation of a cancer cell in a patient which comprises administering to the patient a composition which reduces integrin-integrin interaction, integrin receptor clustering interaction, or integrin-non-integrin protein interaction.
- 90. A method of reducing the proliferation of a cancer cell in a patient which comprises administering to the patient a composition which reduces the production of an integrin or laminin by the cancer cell.
- 91. A method of reducing the proliferation of a cancer cell in a patient which comprises administering to the patient a composition which interfers with an integrin signaling pathway.
- 92. The methods of claims 88-91, wherein the reduction in cancer cell proliferation is selective.
 - 93. The method of any of claims 88-91, wherein the administering is local or systemic.
- 94. The method of any of claims 88-91, wherein the patient is additionally administered an agent which modulates cell signaling, a chemotherapeutic drug, or treated with radiation or angiogenesis inhibitor.
- 95. The method of claim 94, wherein the additional agent is administered serially or in combination.
- 96. The method of claim 93, wherein the local administering is direct application to cancer cells.
- 97. The method of claim 96, wherein the direct application to cancer cells is performed during surgery.
- 98. The method of claim 97, wherein the direct application to cancer cells is performed topically to cancerous tissue.
- 99. The method of claim 94, wherein the systemic administration is by subcutaneous, intraperitoneal, intra-arterial, intravenous, or bolus administration, or by application through a catheter or similar apparatus.
- 100. The method of claim 94, wherein the systemic administration comprises a long-term release formulation.
- 101. The method of claim 94, wherein the systemic administration is by oral administration.
- 102. The method of claim 101 wherein the oral administration comprises administering a pill, capsule, tablet, liquid or suspension.

103. A method of reducing the proliferation of a cancer cell in a patient which comprises administering to a patient a vector comprising coding sequence for a protein or peptide which inhibits ligand binding to integrin receptors on cancer cells wherein the coding sequence is under the control of a promoter which functions in mammalian cells.

- 104. A method of reducing the proliferation of a cancer cell in a patient which comprises administering to a patient a vector comprising coding sequence for a protein or peptide which prevents integrin receptor subunits from interacting with one another, prevents integrin receptor clustering interaction, or prevents integrin receptor subunits from interacting with other proteins on cancer cells wherein the coding sequence is under the control of a promoter which functions in mammalian cells.
- 105. A method of reducing the proliferation of a cancer cell in a patient which comprises administering to a patient a vector comprising coding sequence for a protein or peptide which prevents integrin receptor subunits from interacting with one another, prevents integrin receptor clustering interaction, or prevents integrin receptor subunits from interacting with other proteins in cancer cells wherein the coding sequence is under the control of a promoter which functions in mammalian cells.
- 106. A method of reducing the proliferation of a cancer cell in a patient which comprises administering to a patient a vector comprising coding sequence for a protein or peptide which interferes with integrin subunit or laminin production wherein the coding sequence is under the control of a promoter which functions in mammalian cells.
- 107. A method of reducing the proliferation of a cancer cell in a patient which comprises administering to a patient a vector comprising coding sequence for a protein or peptide which interferes with an integrin signaling pathway of cells wherein the coding sequence is under the control of a promoter which functions in mammalian cells.
- 108. The methods of claims 103-107, wherein the reduction in cancer cell proliferation is selective.
- 109. The method of claims 103-107 wherein the vector is administered directly to a cancer cell.
- 110. The method of claims 103-107, wherein the vector is administered directly to a normal cell.
- 111. The method of claims 103-107, wherein the vector is packaged in a viral vector or liposome.
 - 112. The method of claim 88 wherein the vector is a retroviral vector.

113. The method of any one of claims 103-107, wherein the vector is administered systemically.

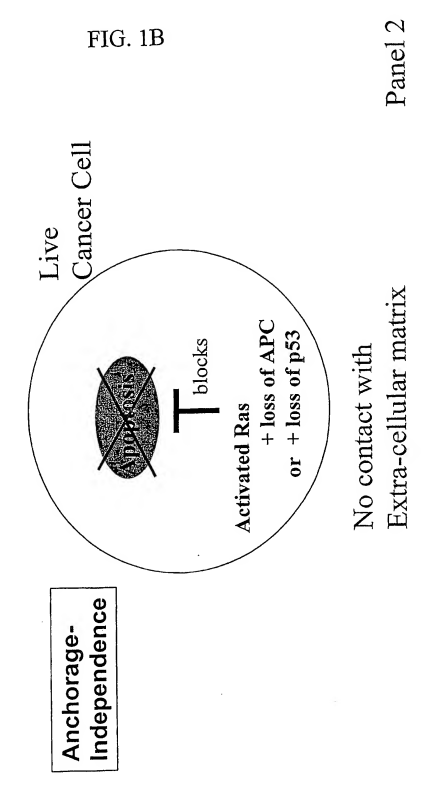
- 114. The method of claim 109, wherein the direct administration is by topical application.
- 115. The method of claim 111, wherein the direct administration is by topical application.
- 116. The method of claim 114, wherein the direct administration is performed during surgery.
- 117. The method of claim 115, wherein the direct administration is performed during surgery.
 - 118. The method of any of claims 103-107, wherein the patient is an animal.
 - 119. The method of claim 118 wherein the animal is a mammal.
 - 120. The method of claim 119, wherein the mammal is human.
- 121. The method of claims 103-107, wherein the cancer cells are selected from the group consisting of melanoma, adenoma, lymphoma, myeloma, carcinoma, plasmocytoma, sarcoma, glioma, thyoma, leukemia, skin cancer, retinal cancer, breast cancer, prostate cancer, colon cancer, esophageal cancer, stomac cancer, pancreas cancer, brain tumors, lung cancer, ovarian cancer, cervical cancer, hepatic cancer, gastrointestinal cancer, and head and neck cancer cells.
- 122. The method of any of claims 103-107 wherein the patient is additionally administered at least one of another agent which modulates cell signaling, a chemotherapeutic drug, an angiogenesis inhibitor or treated with radiation.
- 123. The method of claim 120 wherein the other agent which modifies cell signaling, chemotherapeutic drug, angiogenesis inhibitor or radiation treatment is administered serially or in combination.

Normal cells require integrin-matrix interactions for survival

^{1/8} FIG. 1A

Cell dead Extra-cellular matrix integrin Lost contact with No signal Cell alive Extra-cellular matrix Signal blocks cell death

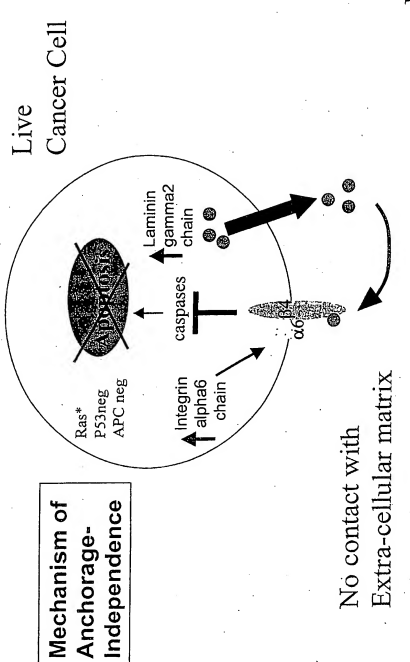
Cells transformed by co-operating oncogenic mutations survive without contact to extra-cellular matrix



Oncogenes co-operate to induce cancer cell survival

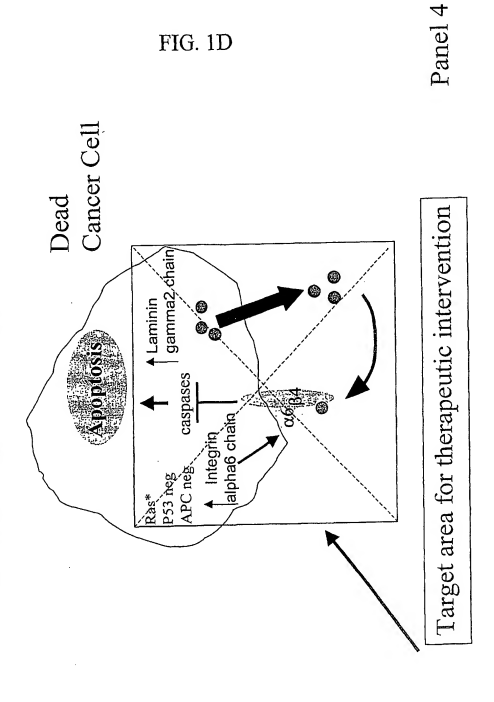
via engagement of autocrine integrin signaling

FIG. 1C

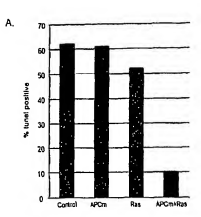


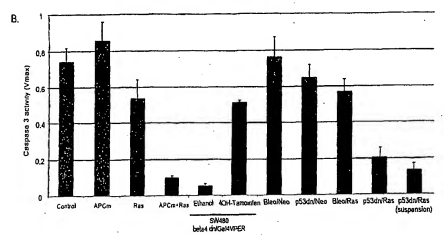
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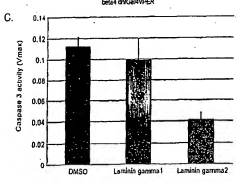
Inhibition of oncogene-induced autocrine integrin signaling loop causes cancer cell death

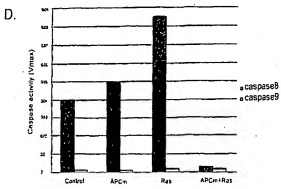


5/8 FIG. 2







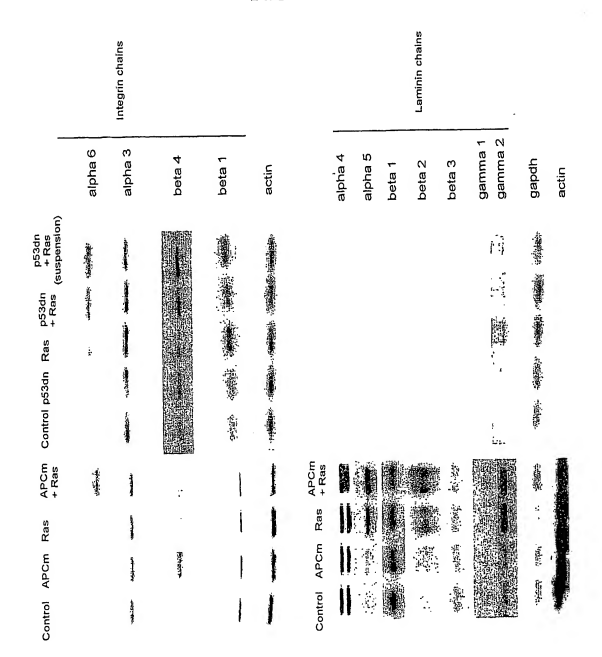


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6/8

FIG. 3



7/8

FIG. 4

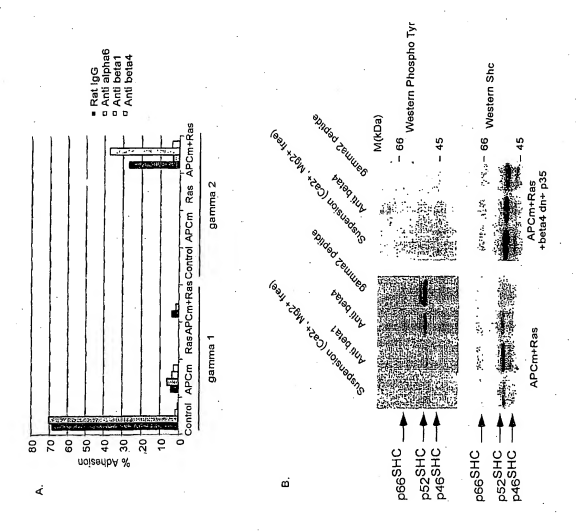
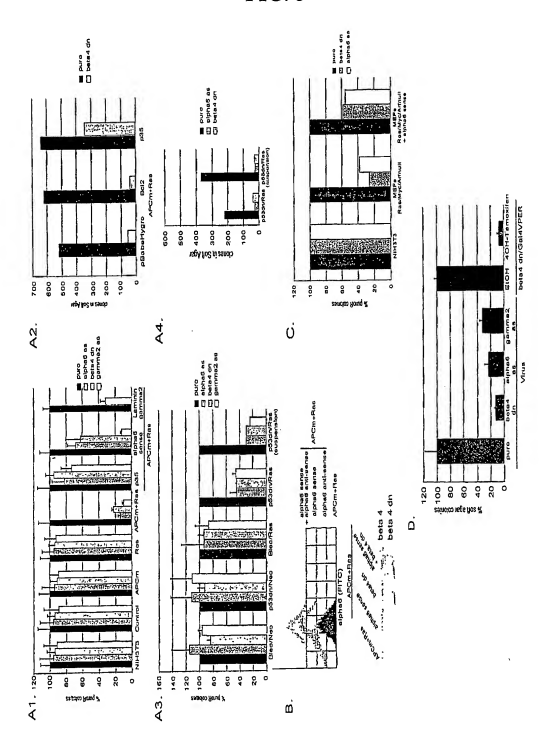


FIG. 5



1

SEQUENCE LISTING

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Deleu, Laurent Land, Hartmut

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<223> Description of Artificial Sequence; Note =
 synthetic construct

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1800
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ccagaagttc ttccaattct gaattcagat gaacccaaga cagctcatat tgatgttcac
                                                                         1860
                                                                         1920
ttcttaaaag agggatgtgg agacgacaat gtatgtaaca gcaaccttaa actagaatat
                                                                         1980
aaattttgca cccgagaagg aaatcaagac aaattttctt atttaccaat tcaaaaaggt
gtaccagaac tagttctaaa agatcagaag gatattgctt tagaaataac agtgacaaac
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                                                                         2280
aatcotttta aaagaaatto aaatgtoact ttttatttgg ttttaagtac aactgaagto
acctttgaca ccccatatct ggatattaat ctgaagttag aaacaacaag caatcaagat
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aatttggctc caattacagc taaagcaaaa gtggttattg aactgctttt atcggtctcg
                                                                         2400
ggagttgcta aaccttccca ggtgtatttt ggaggtacag ttgttggcga gcaagctatg
                                                                         2460
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                                                                         2520
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aatgggaaat ggttgcttta ttiggtgaaa gtagaatcca aaggattgga aaaggtaact
                                                                         2640
tgtgagccac aaaaggagat aaactccctg aacctaacgg agtctcacaa ctcaagaaag
                                                                         2700
                                                                         2760
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agaaaatacc agactcttaa ctgtagcgtg aacgtgaact gtgtgaacat cagatgcccg
                                                                         2820
etgeggggge tggacageaa ggegtetett attitgeget egaggttatg gaacageaca
                                                                         2880
                                                                         2940
tttctagagg aatattccaa actgaactac ttggacattc tcatgcgagc cttcattgat
gtgactgctg ctgccgaaaa tatcaggctg ccaaatgcag gcactcaggt tcgagtgact
                                                                         3000
gtgtttccct caaagactgt agctcagtat tcgggagtac cttggtggat catcctagtg
                                                                         3060
gctattctcg ctgggatctt gatgcttgct ttattagtgt ttatactatg gaagtgtggt ttcttcaaga gaaataagaa agatcattat gatgccacat atcacaaggc tgagatccat
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geteageeat etgataaaga gaggettaet tetgatgeat ag
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<210> 2

<211> 1072

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence; Note =
 synthetic construct

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Thr	Tyr 210	Asn	Trp	Lys	Gly	11e 215	Val	Arg	Val	Glu	Gln 220	Lys	Asn	Asn	Thr
Phe 225		Asp	Met	Asn	Ile 230		Glu	Asp	Gly	Pro 235		Glu	Val	Gly	Gly 240
	Thr	Glu	His	Asp 245		Ser	Leu	Val	Pro 250		Pro	Ala	Asn	Ser 255	
Leu	Gly	Phe	Ser 260		Asp	Ser	Gly	Lуs 265		Ile	Val	Ser	Lys 270	Asp	Glu
Ile	Thr	Phe 275		Ser	Gly	Ala	Pro 280		Ala	Asn	His	Ser 285	-	Ala	Val
Val	Leu 290	_	Lys	Arg	Asp	Met 295		Ser	Ala	Hìs	Leu 300		Pro	Glu	His
Ile 305		Asp	Gly		Gly 310	Leu	Ala	Ser	Ser	Phe 315	Gly	Tyr	Asp	Val	Ala 320
Val	Val	Asp	Leu	Asn 325	Ьуs	Asp	Gly	Trp	Gln 330	Asp	Ile	Val	Ile	Gly 335	Ala
Pro	Gln	Tyr	Phe 340	Asp	Arg	Asp	Gly	Glu 345	Val	Gly	Gly	Ala	Val 350	Tyr	Val
_		355			-	_	360				_	365		Arg	
	370		_	_		375					380	_		Ile	_
385				_	390	-		_		395		_		Pro	400
				405					410					Gly 415	
Asn -		_	420					425					430	Phe	_
		435					440					445		Pro	
	450					455					460			Ser	
465					470				·	475				Arg	480
_		_		485			_	-	490					Cys 495	
			500	-				505					510	Tyr	
•		515					520					525		Arg Ser	
-	530					535				_	540		_	Lys	
54.5					550					555		_		Lys	560
_				565					570			_		575 Ser	
			580					585					590	Asn	
		595					600					605		Glu	
	610					615					620		_	Tyr	
625					630					635				Pro	640
				645					650			_		655 Ile	
		Ile	660					665					670	Pro	
	Asp	675				His	680					685		Phe	
	690					695					700				. •

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Asp Thr Leu Thr Tyr Ser Ala Tyr Arg Glu Leu Arg Ala Phe Pro Glu
                                     715
                  710
Lys Gln Leu Ser Cys Val Ala Asn Gln Asn Gly Ser Gln Ala Asp Cys
                               730
               725
Glu Leu Gly Asn Pro Phe Lys Arg Asn Ser Asn Val Thr Phe Tyr Leu
                                            750
                             745
           740
Val Leu Ser Thr Thr Glu Val Thr Phe Asp Thr Pro Tyr Leu Asp Ile
                      760
                                            765
       755
Asn beu Lys Leu Glu Thr Thr Ser Asn Gln Asp Asn beu Ala Pro Ile
                                         780
                      775
Thr Ala Lys Ala Lys Val Val Ile Glu Leu Leu Ser Val Ser Gly
                                     795
                  790
Val Ala Lys Pro Ser Gln Val Tyr Phe Gly Gly Thr Val Val Gly Glu
                                                    815
                                 810
               805
Gln Ala Met Lys Ser Glu Asp Glu Val Gly Ser Leu Ile Glu Tyr Glu
                                              830
                             825
           820
Phe Arg Val Ile Asn Leu Gly Lys Pro Leu Thr Asn Leu Gly Thr Ala
                                            845
                         840
        835
Thr Leu Asn Ile Gln Trp Pro Lys Glu Ile Ser Asn Gly Lys Trp Leu
                                     860
                      855
   850
Leu Tyr Leu Val Lys Val Glu Ser Lys Gly Leu Glu Lys Val Thr Cys
                   870
                                     875
Glu Pro Gln Lys Glu Ile Asn Ser Leu Asn Leu Thr Glu Ser His Asn
                                                    895
                                 890
               885
Ser Arg Lys Lys Arg Glu Ile Thr Glu Lys Gln Ile Asp Asp Asn Arg
                                                 910
                             905
Lys Phe Ser Leu Phe Ala Glu Arg Lys Tyr Gln Thr Leu Asn Cys Ser
                                            925
                          920
        915
Val Asn Val Asn Cys Val Asn Ile Arg Cys Pro Leu Arg Gly Leu Asp
                      935
                                        940
Ser Lys Ala Ser Leu Ile Leu Arg Ser Arg Leu Trp Asn Ser Thr Phe
                                                        960
                  950
                                     955
Leu Glu Glu Tyr Ser Lys Leu Asn Tyr Leu Asp Ile Leu Met Arg Ala
                                  970
               965
 Phe Ile Asp Val Thr Ala Ala Ala Glu Asn Ile Arg Leu Pro Asn Ala
                                                990
                              985
            980
Gly Thr Gln Val Arg Val Thr Val Phe Pro Ser Lys Thr Val Ala Gln
                                             1005
                          1000
Tyr Ser Gly Val Pro Trp Trp Ile Ile Leu Val Ala Ile Leu Ala Gly
                                       1020
                       1015
    1010
 Ile Leu Met Leu Ala Leu Leu Val Phe Ile Leu Trp Lys Cys Gly Phe
                                     1035
            1030
 Phe Lys Arg Asn Lys Lys Asp His Tyr Asp Ala Thr Tyr His Lys Ala
              1045 1050
                                                    1055
 Glu Ile His Ala Gln Pro Ser Asp Lys Glu Arg Leu Thr Ser Asp Ala
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            1060
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 <211> 3222
 <212> DNA
 <213> Artificial Sequence
 <220>
 <223> Description of Artificial Sequence; Note =
      synthetic construct
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                                                                  120
 gggageetet teggettete getegeeatg caetggeagt tgeageegga ggacaagegg
                                                                   180
                                                                   240
 gggggcctgt acagctgtga catcacctcc cgaggacctt gtacacggat tgaatttgat
                                                                  300
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```
360
aatgacgctg atcctatgtc agaaagcaag gaagaccagt ggatgggagt cactgtccag
agccaaggtc cagggggcaa agtggtgacg tgtgcacatc gatatgagaa acggcagcac
                                                                           420
gtcaacacga agcaggagtc gcgggatatc tttggaagat gttatgtcct gagtcagaat
                                                                           480
                                                                           540
ctcagaattg aagatgatat ggacggagga gactggagtt tctgcgatgg ccggttgaga
ggccatgaaa agtttggctc ctgtcagcaa ggagtagcgg ctactttcac taaggacttt
                                                                           600
                                                                           660
cattacattg tittiggage eccagggact tacaactgga aagggategt eegtgtagaa
caaaagaata acacttitit tgacatgaac atcittgaag atgggcccta tgaagttggt
                                                                           720
                                                                           780
ggagagacag atcatgatga aagtotogtg cocgttootg ctaacagtta cotaggotit
                                                                           840
togotggact cagggaaggg tattgtttct aaagatgaca toacttttgt gtotggtgct
ccaagagcca atcacagtgg ggctgtagtg ttgctaaaaa gagacatgaa gtccgcacat
                                                                           900
                                                                           960
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                                                                          1020
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tttgataggg atggtgaagt cgggggtgca gtttacgtct acattaacca gcaaggcaaa
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                                                                          1140
                                                                          1200
gtgaaaaata taggtgatat taaccaagat ggctatccag atattgctgt tggagctccc
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ccaacacagg ttctcgaggg gacatcgcct tacttcggct attcaatcgc tgggaatatg
                                                                          1320
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                                                                          1500
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                                                                          1560
                                                                          1620
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aaccaaggtt ccgagccaaa gtatactcag gagctgaccc tgaatcggca gaagcagcgg
                                                                          1680
                                                                          17.40
gcgtgcatgg aggagaccct ctggctgcag gagaacatca gagacaagct gcgtcccatc
cccatcacgg cttctgtgga gatccaggag cccacgtctc gccggcgggt gaactcactc
                                                                          1800
                                                                          1860
cccgaagttc ttcccatcct gaattcaaat gaagccaaaa cggtccagac agatgtccac
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                                                                          1980
                                                                          2040
atoccagaat tagtoctaaa agatoagaaa gatatagoto tggaaataac ggtgaccaac
agecettegg atecaaggaa teeeeggaaa gatggegaeg atgeecatga agecaaacte
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                                                                          2160
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                                                                          2640
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tgtgagccac acaatgaaat aaactacctg aagctgaagg agtctcacaa ctcaagaaag
                                                                          2700
                                                                          2760
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                                                                           2820
aqaaaatacc agacteteaa etgeagegte aacgteaggt gtgtgaacat caggtgeeea
ctgcgagggc tggacacgaa ggcctctctc gttctgtgtt ccaggttgtg gaacagcaca tttctagagg aatattccaa actgaactac ttggacattc tcgtgagggc ttccatagat
                                                                           2880
                                                                           2940
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                                                                           3000
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                                                                           3060
actattetta coggattet gatactaget etattagtat tittactata gaagtatage
                                                                           3120
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ttottoaaga gaaataagaa agatoattao gatgocacot atoacaaggo tgagatocat
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<210> 4
<211> 1073
<212> PRT
<213> Artificial Sequence
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<220>
<223> Description of Artificial Sequence; Note =
 synthetic construct

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Ala	Met 50	His	Trp	Gln	Leu	Gln 55	Pro	Glu	Asp	Lys	Arg 60	Leu	Leu	Leu	Val
Gly 65	Ala	Pro	Arg	Ala	Glu 70	Ala	Leu	Pro	Leu	Gln 75	Arg	Ala	Asn	Arg	Thr 80
Gly	Gly	Leu	Tyr	Ser 85	Cys	Asp	Ile	Thr	Ser 90	Arg	Gly	Pro	Cys	Thr 95	Arg
			100					105					110	G1u	
		115					120					125		Lys	
	130					135					140			Thr	
145					150					155				Gln	100
				165					170					Cys 175	
-			180					185					190	Gly	
		195					200					205		Ala	
	210					215					220			Asn	
225					230					235				Val	240
				245					250					Asn 255	
•			260					265					2/0	Lys	
_		275					280					285		Gly	
	290					295					300			Pro	
305					310					315				Asp	320
				325					330	}				11e 335	
Ala			340)				345)				350		
Val		355	;				360)				365)	Ile	
	370)				375	5				386)		Asn	
385	;				390)				395	5			Ala	400
-				405	;				410)				Thr 415 Tyr	
			420)				425	5				430)	
_	_	43	5				44()				445	5	Tyr	
	450)				45	5				46)		Arg	
46	5				476)				47	õ			Aşn	480
				48	5				49	0				/ Ile 495	•
Lei	ı Lys	∍ va.	500		я су:	2 E116	= ቦፗ/	50.		r Mi	a Lry	a EI	510	c Gly	, тйт

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Asn Pro Pro Ile Ser Ile Leu Gly Ile Leu Glu Ala Glu Lys Glu Arg 515 520 Arg Lys Ser Gly Leu Ser Ser Arg Val Gln Phe Arg Asn Gln Gly Ser 535 Glu Pro Lys Tyr Thr Gln Glu Leu Thr Leu Asn Arg Gln Lys Gln Arg 550 555 Ala Cys Met Glu Glu Thr Leu Trp Leu Gln Glu Asn Ile Arg Asp Lys 565 570 Leu Arg Pro Ile Pro Ile Thr Ala Ser Val Glu Ile Gln Glu Pro Thr 580 585 Ser Arg Arg Arg Val Asn Ser Leu Pro Glu Val Leu Pro Ile Leu Asn 600 Ser Asn Glu Ala Lys Thr Val Gln Thr Asp Val His Phe Leu Lys Glu 615 620 Gly Cys Gly Asp Asp Asn Val Cys Asn Ser Asn Leu Lys Leu Glu Tyr 630 Lys Phe Gly Thr Arg Glu Gly Ash Gln Asp Lys Phe Ser Tyr Leu Pro 650 . 645 Ile Gln Lys Gly Ile Pro Glu Leu Val Leu Lys Asp Gln Lys Asp Ile 660 665 670 Ala Leu Glu Ile Thr Val Thr Asn Ser Pro Ser Asp Pro Arg Asn Pro 680. Arg Lys Asp Gly Asp Asp Ala His Glu Ala Lys Leu Ile Ala Thr Phe 695 700 Pro Asp Thr Leu Thr Tyr Ser Ala Tyr Arg Glu Leu Arg Ala Phe Pro 710 715 Glu Lys Gln Leu Ser Cys Val Ala Asn Gln Asn Gly Ser Gln Ala Asp 730 725 Cys Glu Leu Gly Asn Pro Phe Lys Arg Asn Ser Ser Val Thr Phe Tyr
740 745 750 Leu Ile Leu Ser Thr Thr Glu Val Thr Phe Asp Thr Thr Asp Leu Asp 760 Ile Asn Leu Lys Leu Glu Thr Thr Ser Asn Gln Asp Lys Leu Ala Pro 775 780 Ile Thr Ala Lys Ala Lys Val Val Ile Glu Leu Leu Ser Leu Ser 790 795 Gly Val Ala Lys Pro Ser Gln Val Tyr Phe Gly Gly Thr Val Val Gly 810 Glu Gln Ala Met Lys Ser Glu Asp Glu Val Gly Ser Leu Ile Glu Tyr 820 825 Glu Phe Arg Val Ile Asn Leu Gly Lys Pro Leu Lys Asn Leu Gly Thr 840 845 Ala Thr Leu Asn Ile Gln Trp Pro Lys Glu Ile Ser Asn Gly Lys Trp 855 860 Leu Leu Tyr Leu Met Lys Val Glu Ser Lys Gly Leu Glu Gln Ile Val 870 875 Cys Glu Pro His Asn Glu Ile Asn Tyr Leu Lys Leu Lys Glu Ser His 885 890 Asn Ser Arg Lys Lys Arg Glu Leu Pro Glu Lys Gln Ile Asp Asp Ser 900 905 Arg Lys Phe Ser Leu Phe Pro Glu Arg Lys Tyr Gln Thr Leu Asn Cys 915 920 925 Ser Val Asn Val Arg Cys Val Asn Ile Arg Cys Pro Leu Arg Gly Leu 935 Asp Thr Lys Ala Ser Leu Val Leu Cys Ser Arg Leu Trp Asn Ser Thr 955 950 Phe Leu Glu Glu Tyr Ser Lys Leu Asn Tyr Leu Asp Ile Leu Val Arg 965 970 Ala Ser Ile Asp Val Thr Ala Ala Ala Gln Asn Ile Lys Leu Pro His 985 Ala Gly Thr Gln Val Arg Val Thr Val Phe Pro Ser Lys Thr Val Ala

PCT/US01/32127

Gln Tyr Ser Gly Val Ala Trp Trp Ile Ile Leu Leu Ala Val Leu Ala
1010 1015 1020

Gly Ile Leu Met Leu Ala Leu Leu Val Phe Leu Leu Trp Lys Cys Gly
1025 1030 1035 1040

Phe Phe Lys Arg Asn Lys Lys Asp His Tyr Asp Ala Thr Tyr His Lys
1045 1050 1055

Ala Glu Ile His Thr Gln Pro Ser Asp Lys Glu Arg Leu Thr Ser Asp
1060 1065 1070

<210> 5 <211> 5622 <212> DNA <213> Artificial Sequence <220>

<223> Description of Artificial Sequence; Note =
 synthetic construct

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<223> Description of Artificial Sequence; Note = synthetic construct

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465					Asn 470					475					480
				485	Gly				490					495	
			500		Cys			505					510	_	
Gly		515			Gln		520					525			_
Arg	530				Phe	535					540				
545					Cys 550					555					560
				565	Gly				570					575	
			580		Ile			585					590	•	
		595			GJ A		600					605			-
	610				Glu	615					620				
625.					Pro 630					635					640
				645	Arg				650			_		655	_
Trp			660		Glu			665					670		
		675			Thr		680					685			
	690				Thr	695					700				
705					Phe 710					715					720
				. 725	Leu				730		_	-	_	735	
Cys		•	740		Leu			745					750		
	•	755			Glu		760	-			_	765			
	770				Asp	775					780				_
785					Arg 790					795				_	800
				805	Ala	•			810					815	
			820		Arg			825					830		
		835			Glu		840					845			
	850				Tyr	855					860				
865					Gln 870					875					880
				885	Val Glu				890					895	
			900					905					910		
		915			Gly		920					925			
	930				Phe	935					940				-
945		Leu	7116	**	Arg 950	E T.O	GIU	лър	мэр	955	GIU	nλε	GTD	ьеи	Leu 960

Val Glu Ala Ile Asp Val Pro Ala Gly Thr Ala Thr Leu Gly Arg Arg 970 965 Leu Val Asn Ile Thr Ile Ile Lys Glu Gln Ala Arg Asp Val Val Ser 990 985 980 Phe Glu Gln Pro Glu Phe Ser Val Ser Arg Gly Asp Gln Val Ala Arg 1005 995 1000 The Pro Val The Arg Arg Val Leu Asp Gly Gly Lys Ser Gln Val Ser 1010 1015 1020 Tyr Arg Thr Gln Asp Gly Thr Ala Gln Gly Asn Arg Asp Tyr Ile Pro 1025 1030 1035 1040 Val Glu Gly Glu Leu Leu Phe Gln Pro Gly Glu Ala Trp Lys Glu Leu 1045 1050 1055 Gln Val Lys Leu Leu Glu Leu Gln Glu Val Asp Ser Leu Leu Arg Gly 1060 1065 1070 Arg Gln Val Arg Arg Phe His Val Gln Leu Ser Asn Pro Lys Phe Gly 1085 1075 1080 Ala His Leu Gly Gln Pro His Ser Thr Thr Ile Ile Ile Arg Asp Pro 1090 1095 1100 Asp Glu Leu Asp Arg Ser Phe Thr Ser Gln Met Leu Ser Ser Gln Pro 1110 1115 Pro Pro His Gly Asp Leu Gly Ala Pro Gln Asn Pro Asn Ala Lys Ala 1125 1130 1135 Ala Gly Ser Arg Lys Ile His Phe Asn Trp Leu Pro Pro Ser Gly Lys 1140 1145 1150 Pro Met Gly Tyr Arg Val bys Tyr Trp Ile Gln Gly Asp Ser Glu Ser 1155 1160 1165 Glu Ala His Leu Leu Asp Ser Lys Val Pro Ser Val Glu Leu Thr Asn 1170 1175 1180 Leu Tyr Pro Tyr Cys Asp Tyr Glu Met Lys Val Cys Ala Tyr Gly Ala 1190 1195 1200 Gln Gly Glu Gly Pro Tyr Ser Ser Leu Val Ser Cys Arg Thr His Gln 1205 1210 1215 Glu Val Pro Ser Glu Pro Gly Arg Leu Ala Phe Asn Val Val Ser Ser 1220 1225 1230 Thr Val Thr Gln Leu Ser Trp Ala Glu Pro Ala Glu Thr Asn Gly Glu 1235 1240 1245 Ile Thr Ala Tyr Glu Val Cys Tyr Gly Leu Val Asn Asp Asp Asn Arg 1250 1255 1260 Pro Ile Gly Pro Met Lys Lys Val Leu Val Asp Asn Pro Lys Asn Arg 1265 1270 1275 1280 Met Leu Leu Ile Glu Asn Leu Arg Glu Ser Gln Pro Tyr Arg Tyr Thr 1285 1290 1295 Val Lys Ala Arg Asn Gly Ala Gly Trp Gly Pro Glu Arg Glu Ala Ile 1300 1305 1310 Ile Asn Leu Ala Thr Gln Pro Lys Arg Pro Met Ser Ile Pro Ile Ile 1315 1320 1325 Pro Asp Ile Pro Ile Val Asp Ala Gln Ser Gly Glu Asp Tyr Asp Ser 1330 1335 1340 Phe Leu Met Tyr Ser Asp Asp Val Leu Arg Ser Pro Ser Gly Ser Gln 1345 1350 1355 Arg Pro Ser Val Ser Asp Asp Thr Gly Cys Gly Trp Lys Phe Glu Pro 1365 1370 1375 Leu Leu Gly Glu Glu Leu Asp Leu Arg Arg Val Thr Trp Arg Leu Pro 1380 1385 1390 Pro Glu Leu Ile Pro Arg Leu Ser Ala Ser Ser Gly Arg Ser Ser Asp 1395 1400 1405 Ala Glu Ala Pro His Gly Pro Pro Asp Asp Gly Gly Ala Gly Gly Lys 1410 1415 1420 Gly Gly Ser Leu Pro Arg Ser Ala Thr Pro Gly Pro Pro Gly Glu His 1425 1430 1435 1440 Leu Val Asn Gly Arg Met Asp Phe Ala Phe Pro Gly Ser Thr Asn Ser 1445 1450

Leu His Arg Met Thr Thr Thr Ser Ala Ala Ala Tyr Gly Thr His Leu 1460 1465 1470 Ser Pro His Val Pro His Arg Val Leu Ser Thr Ser Ser Thr Leu Thr 1475 1480 1485 Arg Asp Tyr Asn Ser Leu Thr Arg Ser Glu His Ser His Ser Thr Thr 1495 1490 1500 Leu Pro Arg Asp Tyr Ser Thr Leu Thr Ser Val Ser Ser His Gly Leu 1510 1515 Pro Pro Ile Trp Glu His Gly Arg Ser Arg Leu Pro Leu Ser Trp Ala 1525 1530 1535 Leu Gly Ser Arg Ser Arg Ala Gln Met Lys Gly Phe Pro Pro Ser Arg 1540 1545 1550 Gly Pro Arg Asp Ser Ile Ile Leu Ala Gly Arg Pro Ala Ala Pro Ser 1555 1560 1565 Trp Gly Pro Asp Ser Arg Leu Thr Ala Gly Val Pro Asp Thr Pro Thr 1570 1575 1580 Arg Leu Val Phe Ser Ala Leu Gly Pro Thr Ser Leu Arg Val Ser Trp 1590 1595 Gln Glu Pro Arg Cys Glu Arg Pro Leu Gln Gly Tyr Ser Val Glu Tyr 1605 16**1**0 1615 Gln Leu Leu Asn Gly Gly Glu Leu His Arg Leu Asn Ile Pro Asn Pro 1620 1625 Ala Gln Thr Ser Val Val Val Glu Asp Leu Leu Pro Asn His Ser Tyr 1.635 1640 1645 1635 Val Phe Arg Val Arg Ala Gln Ser Gln Glu Gly Trp Gly Arg Glu Arg 1650 1655 1660 Glu Gly Val Ile Thr Ile Glu Ser Gln Val His Pro Gln Ser Pro Leu 1670 1675 Cys Pro Leu Pro Gly Ser Ala Phe Thr Leu Ser Thr Pro Ser Ala Pro 1685 1690 1695 Gly Pro Leu Val Phe Thr Ala Leu Ser Pro Asp Ser Leu Gln Leu Ser 1705 1700 1710 Trp Glu Arg Pro Arg Pro Asn Gly Asp Ile Val Gly Tyr Leu Val 1715 1720 1725 Thr Cys Glu Met Ala Gln Gly Gly Gly Pro Ala Thr Ala Phe Arg Val 1730 1735 1740 Asp Gly Asp Ser Pro Glu Ser Arg Leu Thr Val Pro Gly Leu Ser Glu 1750 1755 1760 Asn Val Pro Tyr Lys Phe Lys Val Gln Ala Arg Thr Thr Glu Gly Phe 1765 1770 1775 Gly Pro Glu Arg Glu Gly Ile Ile Thr Ile Glu Ser Gln Asp Gly Gly 1780 1785 1790 Pro Phe Pro Gln Leu Gly Ser Arg Ala Gly Leu Phe Gln His Pro Leu 1795 1800 1805 Gln Ser Glu Tyr Ser Ser Ile Ser Thr Thr His Thr Ser Ala Thr Glu 1815 1820 Pro Phe Leu Val Gly Pro Thr Leu Gly Ala Gln His Leu Glu Ala Gly 1830 1835 Gly Ser Leu Thr Arg His Val Thr Gln Glu Phe Val Ser Arg Thr Leu 1850 1845 1855 Thr Thr Ser Gly Thr Leu Ser Thr His Met Asp Gln Gln Phe Phe Gln 1860 1865

<210> 7

<211> 5907

<212> DNA

<213> Artificial Sequence

^{:220&}gt;

<223> Description of Artificial Sequence; Note =

synthetic construct

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	- Facecaaaa	: atootooadi	t tecaqqaqq	g catagaget	g gragarara	3180
~~ataccc	+ cttcatccd	, cctdaddato	s acqatqaqaa	a geagerger	g grggaggeea	3240
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	c acaddataai	: acadcacac	a aacaccaaa	a crargeree	c gragaaggaa	3420
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                                                                        5820
gtgctgtcca ggaaccataa agtgggtaga ggtgatacaa ggtctttctg actgcatccc
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Lys Ala Gln Val Lys Ser Cys Thr Glu Cys Ile Arg Val Asp Lys Ser
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                            40
Cys Ala Tyr Cys Thr Asp Glu Leu Phe Lys Glu Arg Arg Cys Asn Thr
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Gln Ala Asp Val Leu Ala Ala Gly Cys Arg Gly Glu Ser Ile Leu Val 70 75 Met Glu Ser Ser Leu Glu Ile Thr Glu Asn Thr Gln Ile Val Thr Ser 90 8.5 Leu His Arg Ser Gln Val Ser Pro Gln Gly Leu Gln Val Arg Leu Arg 110 100 105 Arg Gly Glu Glu Arg Thr Phe Val Phe Gln Val Phe Glu Pro Leu Glu 125 120 115 Ser Pro Val Asp Leu Tyr Ile Leu Met Asp Phe Ser Asn Ser Met Ser 135 140 Asp Asp Leu Asp Asn Leu Lys Gln Met Gly Gln Asn Leu Ala Lys Ile 145 150 160 150 Leu Arg Gln Leu Thr Ser Asp Tyr Thr Ile Gly Phe Gly Lys Phe Val 165 170 Asp Lys Val Ser Val Pro Gln Thr Asp Met Arg Pro Glu Lys Leu Lys 190 185 180 Glu Pro Trp Pro Asn Ser Asp Pro Pro Phe Ser Phe Lys Asn Val Ile 205 200 195 Ser Leu Thr Glu Asn Val Glu Glu Phe Trp Asn Lys Leu Gln Gly Glu 220 210 215 Arg Ile Ser Gly Asn Leu Asp Ala Pro Glu Gly Gly Phe Asp Ala Ile 225 230 240 230 Leu Gln Thr Ala Val Cys Thr Arg Asp Ile Gly Trp Arg Ala Asp Ser 245 250 255Thr His Leu Leu Val Phe Ser Thr Glu Ser Ala Phe His Tyr Glu Ala 265 260 Asp Gly Ala Asn Val Leu Ala Gly Ile Met Asn Arg Asn Asp Glu Lys 280 285 Cys His Leu Asp Ala Ser Gly Ala Tyr Thr Gln Tyr Lys Thr Gln Asp 295 300 Tyr Pro Ser Val Pro Thr Leu Val Arg Leu Leu Ala Lys His Asn Ile 310 315 305 The Pro He Phe Ala Val Thr Asn Tyr Ser Tyr Ser Tyr Tyr Glu Lys 325 330 335 Leu His Lys Tyr Phe Pro Val Ser Ser Leu Gly Val Leu Gln Glu Asp 340 345 350 340 Ser Ser Asn Ile Val Glu Leu Leu Glu Glu Ala Phe Tyr Arg Ile Arg 355 360 Ser Asn Leu Asp Ile Arg Ala Leu Asp Ser Pro Arg Gly Leu Arg Thr 380 375 Glu Val Thr Ser Asp Thr Leu Gln Lys Thr Glu Thr Gly Ser Phe His 395 390 Ile Lys Arg Gly Glu Val Gly Thr Tyr Asn Val His Leu Arg Ala Val 405 410 Glu Asp Ile Asp Gly Thr His Val Cys Gln Leu Ala Lys Glu Asp Gln 420 425 430 Gly Gly Asn Tie His Leu Lys Pro Ser Phe Ser Asp Gly Leu Arg Met 435 Asp Ala Ser Val Ile Cys Asp Val Cys Pro Cys Glu Leu Gln Lys Glu 450 455 Val Arg Ser Ala Arg Cys His Phe Arg Gly Asp Phe Met Cys Gly His 470 475 Cys Val Cys Asn Glu Gly Trp Ser Gly Lys Thr Cys Asn Cys Ser Thr 490 495 485 Gly Ser Leu Ser Asp Thr Gln Pro Cys Leu Arg Glu Gly Glu Asp Lys 500 505 Pro Cys Ser Gly His Gly Glu Cys Gln Cys Gly Arg Cys Val Cys Tyr 525 520 515 Gly Glu Gly Arg Tyr Glu Gly His Phe Cys Glu Tyr Asp Asn Phe Gln 540 535 Cys Pro Arg Thr Ser Gly Phe Leu Cys Asn Asp Arg Gly Arg Cys Ser 550

Met	Gly	Glu	Cys	Val 565	Cys	Glu	Pro	Gly	Trp 570	Thr	Gly	Arg	Ser	Cys 575	Asp
Суѕ	Pro	Leu	Ser 580	Asn	Ala	Thr	Cys	Ile 585		Ser	Asn	Gly	Gly 590	Ile	Cys
Asn	Gly	Arg 595	Gly	Tyr	Cys	Glu	Cys 600	Gly	Arg	Cys	His	Cys 605	Asn	Gln	Gln
Ser	Leu 610	Tyr	Thr	Asp	Thr	Thr 615	Cys	Glu	Ile	Asn	Tyr 620	Ser	Ala	Ile	Leu
Gly 625	Leu	Cys	Glu	Asp	Leu 630	Arg	Ser	Суѕ	Val	Gln 635	Cys	Gln	Ala	Trp	Gly 640
Thr	Gly	Glu	ъуs	Lys. 645	Gly	Arg	Ala	Cys	Asp 650	Asp	Cys	Pro	Phe	Lys 655	Val
Lys	Met	Val	Asp 660	Glu	Leu	Lys	Lys	Glu 665	Glu	Val	Val	Glu	Tyr 670	Cys	Ser
Phe	Arg	Asp 675	Glu	Asp	Asp	Asp	Cys 680	Thr	Tyr	Ser	Tyr	Asn 685	Val	Glu	Gly
	690					695					70.0		_	Lys	_
705					710					715				Ile	720
				725					730					Tyr 735	
	·		740					745					750	Arg	
	-	755					760					765		Asn	
	770					775					780		_	Asn	
785					790					795				Gln	800
	_			805					810					Leu 815	
Pro		,	820					825					830	Asn	
		835	_		_		840	_				845		Val	
	850					855					860			Lys	
865					870	•				875	-			Gln Lys	880
		,		885					890		_			895 Phe	
			900					905					910	Gln	
		915					920					925			Val
	930					935					940				Leu
945					950					955		-		Gly	960
				965					970					975 Val	_
			980					985					990		Ala
		995					1000).				1005	5.		Val
	1010)	-			1015	5				1020)			.Val
1025	5				1030)				1035	· .	,		Lys	1040
		J2 U	U.L.y	1045		₽€U	EHE	uT2	1050		GTU	1111	1.F	1055	

18

PCT/US01/32127

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<223> Description of Artificial Sequence; Note =
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                                                                                120
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egetgeaaca eccaggegga egttetgget geaggetgea ggggagagag cateetggte
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                                                                                420
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                                                                                900
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                                                                              1920
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                                                                              2040
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                                40
Cys Ala Tyr Cys Thr Asp Glu Leu Phe Lys Glu Arg Arg Cys Asn Thr
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                                                  60
Gln Ala Asp Val Leu Ala Ala Gly Cys Arg Gly Glu Ser Ile Leu Val
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                                             75
Met Glu Ser Ser Leu Glu Ile Thr Glu Asn Thr Gln Ile Val Thr Ser
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Leu	His	Arg	Ser 100	Gln	Val	Ser	Pro	Gln 105	Gly	Leu	Gln	Val	Arg 110	Leu	Arg
Arg	Gly	Glu 115	Glu	Arg	Thr	Phe	Val 120	Phe	Gln	Val	Phe	Glu 125	Pro	Leu	Glu
Ser	Pro 130		Asp	Leu	Tyr	Ile 135		Met	Asp	Phe	Ser 140	Asn	Ser	Met	Ser
Asp 145		Leu	Asp	Asn	Leu 150		Gln	Met	Gly	Gln 155	Asn	Leu	Ala	Lys	Ile 160
Leu	Arg	Gln	Leu	Thr 165	Ser	Asp	Tyr	Thr	Ile 170	Gly	Phe	Gly	Lys	Phe 175	Val
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		195					200					205		Val	
	210					215					220			Gly	
225					230					235				Ala	240
				245					250					Asp 255	
			260					265					270	Glu	
_	_	275					280					285		Glu	
	290					295					300			Gln	
305					310					315				Asn	320
				325					330					Glu 335	
			340					345					350	Glu	
		355					360					365		Ile	
	370					375					380			Arg	
385					390					395				Phe Ala	400
				405					410					415 Asp	
			420					425					430	Arg	
-		435					440					445		Lys	
	450					455					460			Gly	
465					470					475				Ser	480
_				485					490					495 Asp	
			500)				505					510		
		515	i				520)				525	•	Phe	
	530					535	i				540)		Cys	
545					550)				555	i				560
			_	565	5				570)				Cys 575	
Cys	Pro	Leu	Ser 580		Ala	Thr	Cys	585 585	Asp	Ser	: Asn	لا الفيارا	590	Ile	cys

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Asn Gly Arg Gly Tyr Cys Glu Cys Gly Arg Cys His Cys Asn Gln Gln
                            600
                                                 605
Ser Leu Tyr Thr Asp Thr Thr Cys Glu Ile Asn Tyr Ser Ala Ile Leu
                        615
                                             620
Gly Leu Cys Glu Asp Leu Arg Ser Cys Val Gln Cys Gln Ala Trp Gly
625
                    630
                                                             640
Thr Gly Glu Lys Lys Gly Arg Ala Cys Asp Asp Cys Pro Phe Lys Val
                645
                                    650
Lys Met Val Asp Glu Leu Lys Lys Glu Glu Val Val Glu Tyr Cys Ser
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                                665
                                                     670
Phe Arg Asp Glu Asp Asp Asp Cys Thr Tyr Ser Tyr Asn Val Glu Gly
                            680
                                                 685
Asp Gly Ser Pro Gly Pro Asn Ser Thr Val Leu Val His Lys Lys Lys
                                            700
    690
                        695
Asp Cys Leu Pro Ala Pro Ser Trp Trp Leu Ile Pro Leu Leu Ile Phe
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Leu Leu Leu Leu Ala Leu Leu Leu Leu Cys Trp Lys
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<220>

<223> Description of Artificial Sequence; Note =
 synthetic construct

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caggecette aggacattet gggagaaget cagattteag aaggggeaat gagageegtt
                                                                        2040
                                                                        2100
gctgtccggc tggccaaggc aaggagccaa gagaacgact acaagacccg cctggatgac
                                                                        2160
ctcaagatga etgcagaaag gateegggee etgggeagte ageateagaa cagagtteag
                                                                         2220
gatacgagca gactcatctc tcagatgcgc ctgagtctgg caggaagcga agctctcttg
gaaaacacta atatecatte ttetgageae tacgtgggge egaatgattt taaaagtetg
                                                                        2280
gctcaggagg ctacaagaaa ggcagacagc cacgctgagt cagctaacgc aatgaagcaa
                                                                        2340
                                                                         2400
ctagcaaggg aaactgagga ctactccaaa caagcacttt cattggcccg caagctcttg
agtggaggag geggaagtgg ctcttgggac agctccgtgg tacaaggtct tatgggaaaa
                                                                         2460
                                                                         2520
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gaagetgata ggtegtatea geacagtete egeeteetgg attetgeete teagetteag
                                                                         2580
                                                                         2640
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ctctcaaacc tggtgaccag acaaacggat gcattcacgc gtgtgcgaaa caatctgggg
                                                                         2700
aactgggaaa aagaaacacg gcagctttta cagactggaa aggataggag acagacttca
                                                                         2760
gatcagetge tttcccgtge caacettget aaaaacagag cecaagaage getaagtatg
                                                                         2820
ggcaatgcca ctttttatga agttgagaac atcctgaaga acctccgaga gtttgatctg
                                                                         2880
caggttgaag acagaaaagc agaggctgaa gaggccatga agagactctc ctctattagc
                                                                         2940
                                                                         3000
cagaaggttg cggatgccag tgacaagacc cagcaagcag aaacggccct ggggagcgcc
actgccgaca cccaacgggc aaagaacgca gctagggagg ccctggagat cagcagcgag
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                                                                         3120
                                                                         3180
agaaaggage tggagtttga cacggataag gacacggtge agetggtgat tactgaagee
                                                                         3240
                                                                         3300
cagcaagetg atgccagage cacgagtgee ggagttacca tecaagacae reteaacaca
ttggacggca tcctacacct catagaccag cctggcagtg tggatgaaga agggatgatg
                                                                         3360
ctattagaac aagggetttt ccaagccaag acccagatca acagtegact teggecettg
                                                                         3420
atgtctgacc tggaggagag ggtgcgtcgg cagaggaacc acctccatct gctggagact
                                                                         3480
agcatagatg gaattettge tgatgtgaag aacctggaga acattegaga caacctgeee
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                                                                         3579
ccaggetget acaataccca agetettgag caacagtga
<210> 12
<211> 1192
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<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence; Note = synthetic construct

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Ser	Ala	Ser 195		His	Ala	Ser	Ala 200		Phe	Ser	Val	His 205		Ile	Thr
Ser	Thr 210	Phe	Ser	Gln	Asp	Val 215	Asp		Trp	Lys	Ala 220	Val	Gln	Arg	Asn
Gly 225	Ala	Pro	Ala	Lys	Leu 230	His	Trp	Ser	Gln	Arg 235	His	Arg	Asp	Val	Phe 240
Ser	Ser	Ala	Arg	Arg 245	Ser	Asp	Pro	Val	Tyr 250		Val	Ala	Pro	Ala 255	Ьys
			260		Gln			265					270		·
		275			Gly		280					285	-		
	290				Leu	295					300				_
305					Gly 310					315			-		320
				325	His				330					335	
			340		Asn			345					350		
		355			Thr		360					365			
	370				Gly	375		•			380				
385					Lys 390					395					400
				405	Ala Glu				410					415	
			420		Asn			425				•	430		
		435			Pro		440					445			
	450				Ser	455			•		460				
465			•		470					475					480
				485	Pro				490					495	
			500		Phe			505					510		
		515			Cys		520					525			
	530				Gln Tyr	535					540		•		•
545					550 Asn					555					560
				565	Glu				570					575	
			580		Phe			585					590		
		595			Cys		600					605			
	610				Gln	615					620				
625					630 Gly					635			_		640 Ile
				645	Ala				650					655	
		Gly	660		Arg			665					670		
		675					680			-		685	-		_

Ser	Gln 690	Glu	Asn	Asp	Tyr	Lys 695	Thr	Arg	Leu	Asp	Asp	Leu	Lys	Met	Thr
Ala 705		Arg	Ile	Arg	Ala 710		Gly	Ser	Gln	His 715	Gln	Asn	Arg	Val	Gln 720
Asp			Arg	725					730					735	
			Leu 740					745					750		
-		755	Asp				760					765			
_	770		Ala			775					780				
785			Tyr		790					795					800
	_	_	Gly	805					810					815	
		_	Lys 820					825					830		
		835	Thr				840					845			
	850		Leu			855					860				
865			Gln		870					875					880
			Leu	885					890					895	
			Gly 900					905					910		
		915	Arg Asn			•	920					925			
	930	_	Val			935					940				
945	_		Asp		950					955					960
			Ser	965					970					975	
			980 Ala					985					990		
		995					100	0				100	5		
	101	0	Leu			101	5				102	0			
102	5		ьуs		103	0				103	5				1040
				104	5				105	0				105	5
			106 Val	0				106	5				107	Ð	
		107	_				108	0				108	5		
	109	0	Ile			109	5				110	0			
110	5				111	0				111	5				1120 Arg
				112	5				113	0				113	
			114	0				114	5				115	0	Asp
		115	5				116	0		-		116	5		Tyr
VOL	117	_	. weu		rigii	117		, wat	, POI	. Jeu	118		- <u>-</u> y	-,-	- , -

PCT/US01/32127

Asn Thr Gln Ala Leu Glu Gln Gln

```
1185
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<211> 3582
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence; Note =
       synthetic construct
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                                                                             120
egggaactic acagacaaac tggtaatgga ticcgetgee teaactgeaa tgacaacact
                                                                             180
gatggcatte actgegagaa gtgcaagaat ggettttace ggcacagaga aagggacege
                                                                             240
tgtttgccct gcaattgtaa ctccaaaggt tctcttagtg ctcgatgtga caactctgga
                                                                             300
cggtgcagct gtaaaccagg tgtgacagga gccagatgcg accgatgtct gccaggcttc cacatgctca cggatgcggg gtgcacccaa gaccagagac tgctagactc caagtgtgac
                                                                             360.
                                                                             420
tqtgacccag ctggcatcgc agggccctgt gacgcgggcc gctgtgtctg caagccagct
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gttactggag aacgctgtga taggtgtcga tcaggttact ataatctgga tggggggaac
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cctgagggct gtacccagtg tttctgctat gggcattcag ccagctgccg cagctctgca
                                                                             600
gaatacagtg tocataagat cacctotace titcatcaag atgitgatgg ciggaagget
                                                                             660
gtccaacgaa atgggtctcc tgcaaagctc caatggtcac agcgccatca agatgtgttt agctcagccc aacgactaga ccctgtctat tttgtggctc ctgccaaatt tcttgggaat
                                                                             720
                                                                             780
caacaggtga gctatgggca aagcctgtcc tttgactacc gtgtggacag aggaggcaga
                                                                            840
cacccatctg cccatgatgt gattctggaa ggtgctggtc tacggatcac agctcccttg
                                                                             900
atgecacttg geaagacact geettgtggg etcaceaaga ettacacatt caggttaaat
                                                                             960
gagcatccaa gcaataattg gagcccccag ctgagttact ttgagtatcg aaggttactg
                                                                            1020
cggaatetea cageceteeg cateegaget acatatggag aatacagtae tgggtacatt gacaatgtga ceetgatte agecegeeet gtetetggag eeccageace etgggttgaa
                                                                            1080
                                                                            1140
cagtgtatat gtcctgttgg gtacaagggg caattctgcc aggattgtgc ttctggctac
                                                                           1200
aagagagatt cagcgagact ggggcctttt ggcacctgta ttccttgtaa ctgtcaaggg
                                                                           1260
ggaggggcct gtgatccaga cacaggagat tgttattcag gggatgagaa tcctgacatt
                                                                            1320
gagtgtgcig actgcccaat tggtttctac aacgatccgc acgacccccg cagctgcaag
                                                                           1380
ccatgtccct gtcataacgg gttcagctgc tcagtgatgc cggagacgga ggaggtggtg
                                                                           1440
tgcaataact gccctcccgg ggtcaccggt gcccgctgtg agctctgtgc tgatggctac
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                                                                           1620
tgtatccaca acacagoogg catctactgo gaccagtgoa aagcaggota cttoggggac
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ccattggctc ccaacccagc agacaagtgt cgagcttgca actgtaaccc catgggctca
                                                                           1740
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aactgtgagc atggagcatt cagctgtcca gcttgctata atcaagtgaa gattcagatg
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gatcagttta tgcagcagct tcagagaatg gaggccctga tttcaaaggc tcagggtggt
                                                                           1920
gatggagtag tacctgatac agagctggaa ggcaggatgc agcaggctga gcaggccctt
                                                                           1980
caggacatto tgagagatgo coagatttoa gaaggtgota goagatocot tggtotocag
                                                                           2040
ttggccaagg tgaggagcca agagaacagc taccagagcc gcctggatga cctcaagatg
                                                                           2100
actgtggaaa gagttcgggc tctgggaagt cagtaccaga accgagttcg ggatactcac
                                                                           2160
aggeteatea eteagatgea getgageetg geagaaagtg aagetteett gggaaacaet aacatteetg ceteagacea etaegtgggg ceaaatgget ttaaaagtet ggeteaggag
                                                                           2220
                                                                           2280
gccacaagat tagcagaaag ccacgttgag tcagccagta acatggagca actgacaagg
                                                                           2340
gaaactgagg actattccaa acaagccctc tcactggtgc gcaaggccct gcatgaagga
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                                                                           2760
gateagetge tttcccgtge caatettget aaaageagag cacaagaage actgagtatg
                                                                           2820
ggcaatgcca ctttttatga agttgagagc atccttaaaa acctcagaga gtttgacctg
                                                                           2880
caggtggaca acagaaaagc agaagctgaa gaagccatga agagactctc ctacatcagc
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3000
cagaaggttt cagatgccag tgacaagacc cagcaagcag aaagagccct ggggagcgct
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                                                                      3060
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                                                                      3120
                                                                      3180
gccatggaaa agggactggc ctctctgaag agtgagatga gggaagtgga aggagagctg
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                                                                      3240
gcccagaagg ttgataccag agccaagaac gctggggtta caatccaaga cacactcaac
                                                                      3300
acattagacg gcctcctgca tctgatggac cagcctctca gtgtagatga agaggggctg
                                                                      3360
                                                                      3420
gtcttactgg agcagaagct ttcccgagcc aagacccaga tcaacagcca actgcggccc
atgatgtcag agctggaaga gagggcacgt cagcagaggg gccacctcca tttgctggag
                                                                      3480
                                                                      3540
acaagcatag atgggattct ggctgatgtg aagaacttgg agaacattag ggacaacctg
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coccagget getacaatac ccaggetett gageaacagt ga
<210> 14
<211> 1193
<212> PRT
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence; Note =
      synthetic construct
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bys Ser Arg Gln Cys Ile Phe Asp Arg Glu Leu His Arg Gln Thr Gly
                             40
Asn Gly Phe Arg Cys Leu Asn Cys Asn Asp Asn Thr Asp Gly Ile His
Cys Glu Lys Cys Lys Asn Gly Phe Tyr Arg His Arg Glu Arg Asp Arg
                                         75
                    70
Cys Leu Pro Cys Asn Cys Asn Ser Lys Gly Ser Leu Ser Ala Arg Cys
                                     90
                85
Asp Asn Ser Gly Arg Cys Ser Cys Lys Pro Gly Val Thr Gly Ala Arg
                                 105
                                                     110
            100
Cys Asp Arg Cys Leu Pro Gly Phe His Met Leu Thr Asp Ala Gly Cys
                                                 125
Thr Gln Asp Gln Arg Leu Leu Asp Ser Lys Cys Asp Cys Asp Pro Ala
                                             140
    130
                         135
Gly Ile Ala Gly Pro Cys Asp Ala Gly Arg Cys Val Cys Lys Pro Ala
                                         155
                     150
Val Thr Gly Glu Arg Cys Asp Arg Cys Arg Ser Gly Tyr Tyr Asn Leu
                                                         175
                                     170
                165
Asp Gly Gly Asn Pro Glu Gly Cys Thr Gln Cys Phe Cys Tyr Gly His
                                                     190
                                 185
            180
Ser Ala Ser Cys Arg Ser Ser Ala Glu Tyr Ser Val His Lys Ile Thr
                                                 205
                             200
Ser Thr Phe His Gln Asp Val Asp Gly Trp Lys Ala Val Gln Arg Asn
                         215
                                             220
    210
Gly Ser Pro Ala Lys Leu Gln Trp Ser Gln Arg His Gln Asp Val Phe
                     230
                                         235
Ser Ser Ala Gln Arg Leu Asp Pro Val Tyr Phe Val Ala Pro Ala Lys
                                     250
                 245
Phe Leu Gly Asn Gln Gln Val Ser Tyr Gly Gln Ser Leu Ser Phe Asp
                                                      270
                                 265
             260
 Tyr Arg Val Asp Arg Gly Gly Arg His Pro Ser Ala His Asp Val Ile
                                                 285
                             280
         275
Leu Glu Gly Ala Gly Leu Arg Ile Thr Ala Pro Leu Met Pro Leu Gly
                         295
                                             300
 Lys Thr Leu Pro Cys Gly Leu Thr Lys Thr Tyr Thr Phe Arg Leu Asn
```

Glu	His	Pro	Ser	Asn 325	Asn	Trp	Ser	Pro	Gln 330	Leu	Ser	Tyr	Phe	Glu 335	Tyr
Arg	Arg	Leu	Leu 340	Arg	Asn	Leu	Thr	Ala 345	Leu	Arg	Ile	Arg	Ala 350	Thr	Tyr
Gly	Glu	Tyr 355	Ser	Thr	Gly	Tyr	Ile 360	Asp	Asn	Val	Thr	Leu 365	Ile	Ser	Ala
Arg	Pro 370	Val.	Ser	GЉУ	Ala	Pro 375	Ala	Pro	Trp	Val	Glu 380	Gln	Cys	Ile	Cys
Pro 385	Val	Gly	Tyr	Lys	Gly 390	Gln	Phe	Cys	Gln	Asp 395	Cys	Ala	Ser	Gly	Tyr 400
Ъуs	Arg	Asp	Ser	Ala 405	Arg	Leu	Gly	Pro	Phe 410	Gly	Thr	Cys	Ile	Pro 415	Cys
Asn	Cys	Gln	Gly 420	GJÀ	Gly	Ala	Cys	Asp 425	Pro	Asp	Thr	Gly	Asp 430	Cys	Tyr
	-	435				-	440		-			445	Pro		
	450					455					460		Cys		
465					470					475			Glu		480
				485				•	490				Glu	495	
			500	•				505	-				Pro 510		
		515					520					525	Ser		
-	530	_	_	-		535					540		Ile		
545					550	•				555			Phe		560
				565				_	570				Asn	575	
		_	580				-	585	•		•		Thr 590		
_	-	595	_				600					605	Ala Gln		
-	610					615					620				Gly
625					630					635			Gln		640
				645					650				Ser	655	
			660					665					670 Ser		
		675					680					685			Arg
	690					695					700				His
705					710					715					720 Ser
_				725					730					735	Asn
			740					745					750		His
		755			•	•	760					765			Asp
	770					775					780				Gly
785					790					795					800 Leu
	-			805	-			-	810					815	

WO 02/30465 PCT/US01/32127

```
Val Glu Lys Leu Glu Lys Thr Lys Ser Leu Ala Gln Gln Leu Thr Arg
                         825
                             830
        820
Glu Ala Thr Gln Ala Glu Ile Glu Ala Asp Arg Ser Tyr Gln His Ser
                                    845
             840
      835
Leu Arg Leu Leu Asp Ser Val Ser Arg Leu Gln Gly Val Ser Asp Gln
 850 855 860
Ser Phe Gln Val Glu Glu Ala Lys Arg Ile Lys Gln Lys Ala Asp Ser
865 870
                             875
Leu Ser Thr Leu Val Thr Arg His Met Asp Glu Phe Lys Arg Thr Gln
                           890 895
           885
Lys Asn Leu Gly Asn Trp Lys Glu Glu Ala Gln Gln Leu Leu Gln Asn
                         905
                                  910
Gly Lys Ser Gly Arg Glu Lys Ser Asp Gln Leu Leu Ser Arg Ala Asn
                                      925
                     920
Leu Ala Lys Ser Arg Ala Gln Glu Ala Leu Ser Met Gly Asn Ala Thr
                 935
                           940
Phe Tyr Glu Val Glu Ser Ile Leu Lys Asn Leu Arg Glu Phe Asp Leu
        950
                        955
Gln Val Asp Asn Arg Lys Ala Glu Ala Glu Glu Ala Met Lys Arg Leu
          965 970 975
Ser Tyr Ile Ser Gln Lys Val Ser Asp Ala Ser Asp Lys Thr Gln Gln
                         985
Ala Glu Arg Ala Leu Gly Ser Ala Ala Ala Asp Ala Gln Arg Ala Lys
                    1000 1005
     995
Asn Gly Ala Gly Glu Ala Leu Glu Ile Ser Ser Glu Ile Glu Gln Glu
  1010 1015
                                 1020
Ile Gly Ser Leu Asn Leu Glu Ala Asn Val Thr Ala Asp Gly Ala Leu
      1030 1035 1040
Ala Met Glu Lys Gly Leu Ala Ser Leu Lys Ser Glu Met Arg Glu Val
1045 1050 1055
Glu Gly Glu Leu Glu Arg Lys Glu Leu Glu Phe Asp Thr Asn Met Asp 1060 1065 1070
Ala Val Gln Met Val Ile Thr Glu Ala Gln Lys Val Asp Thr Arg Ala
                       1080 1085
Lys Asn Ala Gly Val Thr Ile Gln Asp Thr Leu Asn Thr Leu Asp Gly
                 1095 1100
Leu Leu His Leu Met Asp Gln Pro Leu Ser Val Asp Glu Glu Gly Leu
1105 1110 1115
                                                1120
Val Leu Leu Glu Gln Lys Leu Ser Arg Ala Lys Thr Gln Ile Asn Ser
            1125 1130 1135
Gln Leu Arg Pro Met Met Ser Glu Leu Glu Glu Arg Ala Arg Gln Gln
         1140 1145 1150
Arg Gly His Leu His Leu Leu Glu Thr Ser Ile Asp Gly Ile Leu Ala
      1155 1160 1165
Asp Val Lys Asn Leu Glu Asn Ile Arg Asp Asn Leu Pro Pro Gly Cys 1170 1175 1180
Tyr Asn Thr Gln Ala Leu Glu Gln Gln
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1185
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 <223> Description of Artificial Sequence; Note =
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Lys	Ser	Arg 35	Gln	Cys	Ile	Phe	Asp 40	Arg	Glu	Leu	His	Arg 45	Gln	Thr	Gly
Asn	Gly 50	Phe	Arg	Cys	Leu	Asn 55	Cys	Asn	Asp	Asn	Thr 60	Asp	Gly	Ile	His
Cys 65	Glu	Lys	Cys	Lys	Asn 70	Gly	Phe	Tyr	Arg	His 75	Arg	Glu	Arg	Asp	Arg 80
Cys	Leu	Pro	Cys	Asn 85	Cys	Asn	Ser	Ъуs	Gly 90	Ser	Leu	Ser	Ala	Arg 95	Cys
Asp	Asn	Ser	Gly 100	Arg	Cys	Ser	Cys	Lys 105	Pro	Gly	Val	Thr	Gly 110	Ala	Arg
		115	Cys				120					125			
	130		G l n			135					140				
145			Gly	•	150					155			_		160
			Glu	165					170					175	
			Asn 180				5	185					190		
		195	Cys.		-		200		_			.205	_		
	210	•	His			215					220				
225			Ala		230					235					240
			Gln	245					250					255	
			Asn 260					265					270		_
		275	Asp				280		•			285	_		
	290	•	Ala			295					300				_
305			Pro		310					315			_	,	320
•			Ser	325					330					335	_
			Leu 340					345				_	350		_
		355	Ser		_	_	360	_				365			
	370		Ser	-		375					380		-		_
385			Tyr		390					395					400
			Ser Gly	405					410			_		415	_
			420					425					430		
	•	435	Glu Asp				440					445			-
	450					455					460				
465			Phe		470					475					480
			Cys	485				,	490					495	-
TTC	vañ	O.T.y	Tyr 500	FIIG	атЪ	ASP	ETO	505	атЪ	OTD	מדני	стÀ	510	val	Arg

Pro	Cys	Gln 515	Pro	Cys	Gln	Cys	Asn 520	Asn	Asn	Val	Asp	Pro 525	Ser	Ala	Ser
Gly	Asn 530	Cys	Asp	Arg	Leu	Thr 535	Gly	Arg	Cys	Leu	Lys 540	Cys	Ile	His	Asn
Thr 545	Ala	Gly	Ile	Tyr	Cys 550	Asp	Gln	Cys	Lys	Ala 555	Gly	Tyr	Phe	Gly	Asp 560
Pro	Leu	Ala	Pro	Asn 565	Pro	Ala	Asp	Lys	Cys 570	Arg	Ala	Cys	Asn	Cys 575	Asn
Pro	Met	Gly	Ser 580	Glu	Pro	Val	Gly	Cys 585	Arg	Ser	Asp	Gly	Thr 590	Cys	Val
Cys	Lys	Pro 595	Gly	Phe	Gly	Gly	Pro 600	Asn	Cys	Glu	His	Gly 605	Ala	Phe	Ser
	610		Cys		•	615		_			620				
625			Gln		630					635					640
	_		Val	645					650	-				655	
			Leu 660					665					670		
		675	Ser		_		680			_		685			
	690		Gln Leu			695					700				
705			Thr		710					715					720
			Thr	725					730					735	
	_		740 Ser					745	_				750		
_		755	Ala				760					765			
	770		Gln			775					780				-
785			Gly		790					795					800
			Leu	805					810					815	
			820 Gln		_			825					830		
Leu		835 Leu	Leu	Asp	Ser	Val	840 Ser	Arg	Leu	Gln	Gly	845 Val	Ser	Asp	Gln
Ser	850 Phe	Gln	Val	Glu	G l u	855 Ala	Lys	Arg	Ile	Lys	860 Gln	Lys	Ala	Asp	Ser
865 Le u	Ser	Thr	Leu		870 Thr	Arg	His	Met		875 Gl u	Phe	Lys	Arg		880 Gln
Lys	Asn	Leu	Gly	885 Asn	Trp	Lys	Glu		890 Ala	Gln	Gln	Leu		895 Gln	Asn
Gly	Lys	Ser 915	900 Gly	Arg	Glu	Lys	Ser 920		Gln	Leu	Leu	Ser 925		Ala	Asn
Leu	Ala 930		Ser	Arg	Ala	G1n 935			Leu	Ser	Met 940			Ala	Thr
Phe 945		Glu	Val	Glu	Ser 950		Leu	Ьys	Asn	Leu 955		Glu	Phe	Asp	Leu 960
_	Val	Asp	Asn	Arg 965		Ala	Glu	Ala	Glu 970		Ala	Met	Lys	Arg 975	
Ser	Tyr	Ile	Ser 980		Lys	Val	Ser	Asp 985		Ser	Asp	Lys	Thr 990		Gln
Ala	Glu	Arg 995	Ala	Leu	Gly	Ser	Ala 100		Ala	Asp	Ala	Gln 100	_	Ala	Lys

PCT/US01/32127

2220

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Asn Gly Ala Gly Glu Ala Leu Glu Ile Ser Ser Glu Ile Glu Gln Glu
    1010
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Ile Gly Ser Leu Asn Leu Glu Ala Asn Val Thr Ala Asp Gly Ala Leu
1025
                      1030
                                            1035
Ala Met Glu Lys Gly Leu Ala Ser Leu Lys Ser Glu Met Arg Glu Val
                 1045
                                        1050
Glu Gly Glu Leu Glu Arg Lys Glu Leu Glu Phe Asp Thr Asn Met Asp
             1060
                                   1065
Ala Val Gln Met Val Ile Thr Glu Ala Gln Lys Val Asp Thr Arg Ala
                              1080
                                                     1085
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<223> Description of Artificial Sequence; Note =
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gtccgtgtgg ataaggactg cgcctactgc acagacgaga tgttcaggga ccggcgctgc
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aacacccagg cggagctgct ggccgcgggc tgccagcggg agagcatcgt ggtcatggag
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agcagettee aaateacaga ggagaceeag attgacaeea eeetgeggeg eagceagatg
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tccccccaag gcctgcgggt ccgtctgcgg cccggtgagg agcggcattt tgagctggag
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gtgtttgagc cactggagag ccccgtggac ctgtacatcc tcatggactt ctccaactcc
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atgtccgatg atctggacaa cctcaagaag atggggcaga acctggctcg ggtcctgagc
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                                                                             780
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                                                                             960
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                                                                            1020
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                                                                            1140
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cggggggaag tgggtatata ccaggtgcag ctgcgggccc ttgagcacgt ggatgggacg
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                                                                            1500
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                                                                            1560
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                                                                            1620
cagtgtcccc gcacttccgg gttcctctgc aatgaccgag gacgctgctc catgggccag .
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                                                                            1740
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                                                                            1800
                                                                            1860
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                                                                            1920
agaagaaggg gcgcacgtgt gaggaatgca acttcaaggt caagatggtg gacgagctta
                                                                            1980
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                                                                            2040
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                                                                            2100
agaagaaggg actgecetee gggeteette tggtggetea teeccetget ceteete etgeegetee tggecetget actgetgeta tgetggaagt actgigeetg etgeaagge
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<212> PRT <213> Artificial Sequence

<220>

<223> Description of Artificial Sequence; Note =
 synthetic construct

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Ile	His	Leu 435	Lys	Pro	Ser	Phe	Ser 440	Asp	Gly	Leu	Lys	Met 445	Asp	Ala	Gly
Ile	Ile 450		Asp	Val	Cys	Thr 455		Glu	Leu	Gln	Ьуs 460		Val	Arg	Ser
Ala 465		Cys	Ser	Phe	Asn 470		Asp	Phe	Val	Cys 475	Gly	Gln	Cys	Val	Cys 480
				485					490					Ser 495	
			500					505					510	Cys	
		515					520					525		Gl u	
	530					535					540			Pro	
545					550					555				Gly	560
				565					570					Pro 575	
			580					585					590	Gly	
		595					60.0					605		Leu	
	610					615					620			Ala	
625	_				630					635				Pro	640
				645					650					Arg 655 Pro	
-			660	_				665					670	Val	
-		675					680					685		Arg	
	690					695					700			Leu	
705					710					715				Cys	720
				725					730					735 Gly	
_	_		740					745					750		
		755					760					765		Leu	
	770	_				775					780			Arg	
785	-	_			790					795				Val	800
_				805					810					815	
_			820					825					830		Glu
-		835					840					845)		Gln
	850					855					860				His
865		-			870					875					880 Ala
				885					890	1				895	
		_	900	•				905					910)	Ala
ກະບ	. nys	915			~-y	~]	920					925	5		

Arg	Gly 930	Met	Val	Glu	Phe	Gln 935	Glu	Gly	Val	Glu	Leu 940	Val	Asp	Val	Arg
Val 945			Phe	Ile	Arg 950	Pro	Glu	Asp	Asp	Asp 955	Glu	ГÀ2	Gln	Leu	Leu 960
		Ala	Ile	Asp 965		Pro	Ala	Gly	Thr 970	Ala	Thr	Leu	Gly	Arg 975	Arg
Leu	Val	Asn	Ile 980		Ile	Ile	Lys	Glu 985		Ala	Arg	Asp	Val 990	Val	Ser
Phe	Glu	Gln 995		Glu	Phe	Ser	Val 1000	Ser	Arg	Gly	Asp	Gln 1005	Va1	Ala	Arg
Ile	Pro	Val	Ile	Arg	Arg	Val 1015	Leu		Gly	Gly	Lys 1020	Ser		Val	Ser
Tyr 1025	Arg		Gln	Asp	Gly 1030	Thr		Gln	Gly	Asn 103	Arg		Tyr	Ile	Pro 1040
		GJÀ.	Glu	Leu 1045	Leu		Gln	Pro	Gly 1056	Glu		Trp	Lys	Glu 1055	Leu
Gln	Val	Lys	Leu 1060	Leu		Leu	Gln	Glu 1065	Val		Ser	Leu	Leu 1070	Arg	
Arg	Gln	Val 1075	Arg		Phe	His	Val 1080	Gln		Ser	Asn	Pro 1085	Lys	Phe	Gly
Ala	His 1090	Leu		Gln	Pro	His 1095	Ser		Thr	Ile	Ile 1100	Ile		Asp	Pro
Asp 1105	Glu		Asp	Arg	Ser 1110	Phe		Ser	Gl n	Met 1115	Leu		Ser	Gln	Pro 1120
		His	Gly	Asp 1125	Leu		Ala	Pro	Gln 1130	Asn		Asn	Ala	Lys 1135	Ala
Ala	Gly	Ser	Arg 1140	Lys		His	Phe	Asn 1145	Trp		Pro	Pro	Ser 1150	Gly	
Pro	Met	Gly 1155	Tyr		Val	Lys	Tyr 1160	Trp		Gln	Gly	Asp 1169	Ser	G1u	Ser
Glu	Ala 1170	His		Leu	Asp	Ser 1175	Lys		Pro	Ser	Val 1180		Leu	Thr	Asn
Leu 1185	Tyr		Tyr	Cys	Asp 1190		Glu	Met	Lys	Val 119		Ala	Tyr	Gly	Ala 1200
		Glu	Gly	Pro 1205		Ser	Ser	Leu	Val 1210		Cys	Arg	Thr	His 1215	
Glu	Val	Pro	Ser 1220		Pro	Gly	Arg	Leu 1225		Phe	Asn	Val	Val 1230	Ser	Ser
		1235	5				1240)				1245	5	G l y	
	1250).	_			1255	5				1260)		Asn	
1265	; }	-			1270)				127	5			Asn	1280
				1285	5		_		1290)				Tyr 1295	5
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(19) World Intellectual Property Organization International Bureau



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(43) International Publication Date 18 April 2002 (18.04.2002)

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(10) International Publication Number WO 02/030465 A3

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- (25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data:

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- (72) Inventors; and
- (75) Inventors/Applicants (for US only): LAND, Hartmut [US/US]; 195 Hollywood Avenue, Rochester, NY 14618 (US). DELEU, Laurent [US/US]; 104 Westland Avenue, Rochester, NY 14618 (US).

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- (81) Designated States (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW.
- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

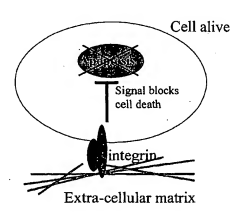
Published:

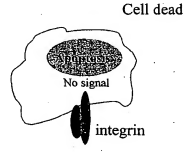
- with international search report
- (88) Date of publication of the international search report: 13 March 2003

[Continued on next page]

(54) Title: COMPOSITIONS THAT INHIBIT PROLIFERATION OF CANCER CELLS

Normal cells require integrin-matrix interactions for survival





Lost contact with Extra-cellular matrix

Panel 1

(57) Abstract: Disclosed are compositions and methods for reducing the proliferation of cancer cells through targeted interactions with integrins.



O 02/030465 A3



For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

Inte Inte Application No PCT/US 01/32127

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 CO7K14/705 A61K A61K38/17 A61K31/7088 A61K38/39 G01N33/566 //A61P35/00 G01N33/50 C12Q1/68 According to International Patent Classification (IPC) or to both national classification and IPC Minimum documentation searched (classification system followed by classification symbols) A61K C12N C07K IPC 7 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) EPO-Internal, BIOSIS, MEDLINE C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Citation of document, with Indication, where appropriate, of the relevant passages TAGLIABUE E ET AL: "Prognostic value of 1-128 X alpha 6 beta 4 integrin expression in breast carcinomas is affected by laminin production from tumor cells." CLINICAL CANCER RESEARCH: AN OFFICIAL JOURNAL OF THE AMERICAN ASSOCIATION FOR CANCER RESEARCH. UNITED STATES FEB 1998, vol. 4, no. 2, February 1998 (1998-02), pages 407-410, XP001119072 ISSN: 1078-0432 abstract introduction page 407 discussion page 409 -page 410 Patent family members are listed in annex. Further documents are listed in the continuation of box C. Special categories of cited documents: *T* later document published after the International filing date or priority date and not in conflict with the application but clied to understand the principle or theory underlying the *A* document defining the general state of the art which is not considered to be of particular relevance "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "E" earlier document but published on or after the international *L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled O document referring to an oral disclosure, use, exhibition or other means in the art. document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of mailing of the International search report Date of the actual completion of the international search 11/12/2002 25 November 2002 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV RIJSWIJK
Tel. (+31-70) 340-2040, Tx. 31 651 epo nl,
Fax: (+31-70) 340-3016 Dumont, E

Inti nel Application No
PCT/US 01/32127

		PC1/US 01/3212/
C.(Continua	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
х	WO 96 10646 A (PYKE CHARLES ;TRYGGVASON KARL (FI); KALLUNKI PEKKA (US)) 11 April 1996 (1996-04-11)	1,7,12, 13,17, 30-34, 40,47, 53,55, 63-65, 89,90
	abstract; claims 1-40	
X	FIELDS ET AL: "Integrins: cell adhesion molecules in cancer" EXPERT OPINION ON THERAPEUTIC PATENTS, ASHLEY PUBLICATIONS, GB, vol. 8, no. 6, August 1998 (1998–08), pages 633-644, XP002105988 ISSN: 1354-3776 abstract Introduction	1,13,24, 30,34
X	NICKOLS A ET AL: "ANTIANGIOGENIC AND ANTICANCER ACTIVITIES OF ANTAGONISTS OF INTEGRINALPHAVBETA3" PROCEEDINGS OF THE 88TH. ANNUAL MEETING OF THE AMERICAN ASSOCIATION FOR CANCER RESEARCH. SAN DIEGO, APR. 12 - 16, 1997, PROCEEDINGS OF THE ANNUAL MEETING OF THE AMERICAN ASSOCIATION FOR CANCER RESEARCH, PHILADELPHIA, AACR, US, vol. 38, 1 March 1997 (1997-03-01), page 206 XP002071773 the whole document	1,13,34
X	WO 98 46265 A (MEYER DEBRA M ;SEARLE & CO (US); NICKOLS G ALLEN (US); SMITH JEFFR) 22 October 1998 (1998-10-22) abstract; claims 1-17	34
A	BACHELDER ROBIN E ET AL: "p53 Inhibits alpha6beta4 integrin survival signaling by promoting the caspase 3-dependent cleavage of AKT/PKB." JOURNAL OF CELL BIOLOGY, vol. 147, no. 5, 29 November 1999 (1999-11-29), pages 1063-1072, XP002222121 ISSN: 0021-9525 abstract page 1070, paragraph 2	1-128
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Intermediate Inter

	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	RABINOVITZ ISAAC ET AL: "The integrin alpha-6-beta-4 and the biology of carcinoma." BIOCHEMISTRY AND CELL BIOLOGY, vol. 74, no. 6, 1996, pages 811-821, XP001120471 ISSN: 0829-8211 the whole document	1-128
T	MERCURIO ARTHUR M ET AL: "Towards a mechanistic understanding of tumor invasion: Lessons from the alpha6beta4 integrin." SEMINARS IN CANCER BIOLOGY, vol. 11, no. 2, April 2001 (2001-04), pages 129-141, XP002222122 ISSN: 1044-579X the whole document especially Table 1, Fig. 3 page 137, column 1, last paragraph -column 2, paragraph 1	1-128
A	D'ABACO GIOVANNA M ET AL: "Synergy between Apc-min and an activated ras mutation is sufficient to induce colon carcinomas." MOLECULAR AND CELLULAR BIOLOGY, vol. 16, no. 3, 1996, pages 884-891, XP002222123 ISSN: 0270-7306 cited in the application the whole document	1-128
A	O'CONNOR KATHLEEN L ET AL: "Release of cAMP gating by the alpha6beta4 integrin stimulates lamellae formation and the chemotactic migration of invasive carcinoma cells." JOURNAL OF CELL BIOLOGY, vol. 143, no. 6, 14 December 1998 (1998-12-14), pages 1749-1760, XP002919093 ISSN: 0021-9525 the whole document	49
Α	CLARKE ASTRID S ET AL: "Activation of the p21 pathway of growth arrest and apoptosis by the beta-4 integrin cytoplasmic domain." JOURNAL OF BIOLOGICAL CHEMISTRY, vol. 270, no. 39, 1995, pages 22673-22676, XP002222124 ISSN: 0021-9258 the whole document	51

Form PCT/ISA/210 (continuation of second sheet) (July 1992)

.....national application No. PCT/US 01/32127

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)	
This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:	
1. X Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely: see FURTHER INFORMATION sheet PCT/ISA/210	
2. X Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically: see FURTHER INFORMATION sheet PCT/ISA/210	
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).	
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)	
This International Searching Authority found multiple inventions in this international application, as follows:	
As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.	
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.	
As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:	
4. No required additional search fees were timely paid by the applicant. Consequently, this international Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:	
Remark on Protest The additional search fees were accompanied by the applicant's protest No protest accompanied the payment of additional search fees.	st.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Continuation of Box I.1

Although claims 1-62, 65-72, as far as they are directed to in vivo methods, and claims 64, 76-128 are directed to a method of treatment of the human/animal body or to a diagnostic method practised on the human/animal body, the search has been carried out and based on the alleged effects of the compounds.

Continuation of Box I.1

Rule 39.1(iv) PCT - Diagnostic method practised on the human or animal body

- Method for treatment of the human or animal body by therapy

Continuation of Box I.2

Present claims 9-12, 14-17, 37-40 relate to a method involving a composition defined by reference to a desirable characteristic or property, namely inhibition of ligand binding to an integrin receptor. The claims cover methods involving all compositions having this characteristic or property, whereas the application provides support within the meaning of Article 6 PCT and disclosure within the meaning of Article 5 PCT for only a limited number of such compositions. In the present case, the claims so lack support, and the application so lacks disclosure, that a meaningful search over the whole of the claimed scope is impossible. Independent of the above reasoning, the claims also lack clarity (Article 6 PCT). An attempt is made to define the composition by reference to a result to be achieved. Again, this lack of clarity in the present case is such as to render a meaningful search over the whole of the claimed scope impossible. Consequently, the search has been carried out for those parts of the claims which appear to be clear, supported and disclosed, namely those parts relating to compositions including antibodies, antisense molecules, functional nucleic acids, primers and probes as defined in the description.

The above reasoning applies by analogy to claims 41, 42 and 62, which relate to methods involving a molecule or an agent defined by reference to a desirable characteristic, i.e. interference with at least one of the cellular proteins mentioned in claim 41, interference with post-translational modifications of integrins, or modulation of cell signaling. The application does not provide support within the meaning of Article 6 PCT and disclosure within the meaning Article 5 PCT for such a molecule or agent. Consequently, a search has not been carried out for those parts of claims 41, 42 and 62 relating to such a molecule or agent.

The applicant's attention is drawn to the fact that claims, or parts of claims, relating to inventions in respect of which no international search report has been established need not be the subject of an

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210 international preliminary examination (Rule 66.1(e) PCT). The applicant is advised that the EPO policy when acting as an International Preliminary Examination Authority is normally not to carry out a preliminary examination on matter which has not been searched. This is the case irrespective of whether or not the claims are amended following receipt of the search report or during any Chapter II procedure.

iformation on patent family members

Inte onal Application No PCT/US 01/32127

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